INTRODUCTION

Transitional cell carcinoma (TCC) urinary bladder is one of the most common as well as lethal urological cancers worldwide accounting for about 95% of bladder cancers (BC).\(^1\) An estimated 81,190 new cases of BC with 17,240 deaths occurred in United States in 2018 and disease is emerging as global menace due to steadily rising life expectancy vis-a-vis heavily incurred health care expenditure (£55 million per year to investigate hematuria in UK).\(^2\) BC incidence and mortality is highest amongst developed nations but lowest in Asia, Latin America and the Caribbean; the variegated geographical distribution being attributed to global variation in risk factors mainly industrialization and smoking exposure.\(^3\)

ABSTRACT

Objective: To determine diagnostic accuracy of NMP 22 and urine cytology in the detection of transitional cell carcinoma (TCC) urinary bladder taking cystoscopy as a gold standard in patients having provisional diagnosis of bladder cancer (BC).

Methods: This cross sectional validational study enrolled 380 patients fulfilling selection criteria and was conducted at Armed Forces Institute of Urology (AFIU) Rawalpindi, Pakistan form July 2018 to July 2019. The urine sample collected underwent NMP22 and cytological analysis followed by rigid cystoscopy. Reports of all three tests divided patients into positive or negative for malignancy as per defined criteria. Sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV) and diagnostic accuracy of NMP 22, urine cytology and their combination was determined. Receiver operating characteristic (ROC) curve analysis performed and area under the curve (AUC) compared among these tests.

Results: The average age of patients was 53.08 ± 12.41 years having male to female ratio 3.75:1(300 males and 80 females). NMP 22 had better sensitivity and comparable specificity to cytology (81.9 & 81.2% vs 54 & 93.9%). Combination of NMP 22 / cytology outperformed both in terms of sensitivity (91.63 vs 81.83 vs 53.96), NPV (87.59 vs 77.46 vs 61.02) and diagnostic accuracy (85.26 vs 81.58 vs 71.32) but at the cost of specificity (76.97 vs 81.21 vs 93.94) and PPV (83.83 vs 85.02 vs 92.06). ROC curve revealed statistically significant higher AUC (0.843 vs .815 vs .73) for combination as compared to NMP 22 and Cytology (p<0.001).

Conclusion: NMP22 is a quick, point of care test having higher sensitivity, NPV and accuracy but similar specificity and PPV to urine cytology for detection of TCC urinary bladder. Combination outperformed both in terms of sensitivity while having modest specificity.

KEYWORDS: Cystoscopy, Nuclear matrix protein 22, Nuclear matrix-associated proteins, Sensitivity, Specificity, Transitional cell carcinoma, Urinary bladder neoplasms, Urine cytology.

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BC carcinogenesis revolves around genetic susceptibility, environmental exposure and unhealthy lifestyles. Majority are non-muscle invasive (NMIBC) having high chances of recurrence as well as progression necessitating lifelong surveillance. Comprehensive, standardized and risk-adapted follow-up protocols incorporating regular cystoscopies, urinary cytology as well as regular upper urinary tract imaging (for high-risk tumors) are recommended by various international bodies like European Association of Urology (EAU), the American Urological association (AUA), society of urological oncology and National comprehensive cancer network (NCCN). In addition to diagnosis, surveillance of BC recurrence pivots cystoscopy as criterion standard due to its high diagnostic accuracy. However, it is invasive, expensive and resource intensive compounded by low patient acceptance as well as poor compliance. Urine cytology is another commonly used gold standard modality for BC screening having excellent specificity. The drawbacks include low sensitivity especially for low grade tumors, requirement of trained cytopathologist and the impossibility of quantification due to subjective interpretation criteria. These compelling limitations lead to approval of several noninvasive biomarkers though with conflicting diagnostic accuracy quoted for primary and recurrent BC.

Nuclear matrix protein (NMP), first described in 1974, is a nonchromatin structure that supports nuclear shape and organizes DNA having up to 25 times higher urinary levels in BC patients. The NMP-22 BladderChek test is an in vitro enzyme immunoassay intended for the qualitative detection of nuclear mitotic apparatus protein (NuMA), an abundant component of NMP in freshly voided spot urine. This painless noninvasive assay provides absolute positive or negative results like pregnancy test within 30 minutes without having any dependency upon intact cancer cells, expert cytopathologist and dedicated laboratory. Low cost, rapid results and simplicity of execution lead to its FDA approval as the only in-office test for the diagnosis of BC.

Although NMP-22 is considered an effective alternative alone or in combination with urine cytology in the diagnosis, screening and surveillance of BC, the clinical evidence generated by several randomized, double-blinded trials conducted recently revealed inconsistent, fluctuating results regarding its diagnostic accuracy. Moreover, to our knowledge very limited local data was found addressing this aspect leading to weak endorsement of use of this potentially-effective diagnostic modality in our setup. We aimed to evaluate the diagnostic accuracy of NMP-22 and voided urine cytology in the detection of TCC in our target population.

METHODS

This cross sectional validation study was conducted at Armed Forces Institute of Urology (AFIU) Rawalpindi, Pakistan form July 2018 to July 2019 after approval of the study protocol from the hospital ethical review board (ERB). Non probability consecutive sampling technique was used to enroll 380 patients after satisfying inclusion criteria (all patients having age 18-75 years of either gender with provisional diagnosis of TCC bladder who were planned to undergo Cystoscopy) while patients already diagnosed with TCC bladder or upper urinary tract, renal malignancy or on dialysis, active gross hematuria, recent history (within two weeks) of urethral instrumentation / catheterization, bladder stones or having active urinary tract infection were excluded. Written informed consent was obtained and demographic details (name, age, and gender) noted.

Voided midstream urine (MSU) was collected at outpatient visit and prior to any treatment, according to the standard protocol for urine collection defined by the Human Kidney and Urine Proteome Project (HKUPP) network. Collected urine was divided into two aliquots; one underwent NMP22 analysis (Qualitative ALERE™ NMP22® BladderChek®) at urology laboratory AFIU according to manufacturers ‘protocol while other portion sent to cytopathology laboratory AFIP, Rawalpindi.

Both air dried and wet fixed slides (ThinPrep slides Cytc Corporation, Marlborough, MA) were prepared by method of direct smearing and centrifuging. After samples were centrifuged at 2000 rpm for five minutes the supernatant was removed to produce cell pellets which were washed with Cytolyt® Solution. Two to three drops of each patient sample was transferred into PreservCyt® Solution and fixed for 15 minutes. Air dried slides were stained with Diff-Quik® stain (MICROPTIC S.L. Barcelona Spain) while Papanicolaou stain used for wet fixed slides. A consultant cytopathologist evaluated all specimens according to Paris classification system for reporting urine cytology. While classes 1 and 2 were considered negative, classes 4 and 5 were deemed positive for bladder cancer, all patients with class 3 (atypical urothelial cells) cytopathological findings but negative cystoscopy were excluded. Presence or absence of urothelial bladder cancer was confirmed with
Rigid cystoscopy under spinal / general anesthesia conducted by consultant urologist having at least 10 years of experience assisted by researcher. All this information was recorded on a specially designed proforma by researcher himself.

**Statistical analysis:** It was performed using IBM SPSS Statistics for Windows, Version 24.0. (Armonk, NY: IBM Corp). Descriptive statistics were used to calculate means ± standard deviation (SD) for quantitative variables e.g., age. The qualitative variable i.e., gender and TCC (on NMP22, urine cytology and cystoscopy) were presented as frequency and percentage and chi square test applied to determine significance. Contingency tables were generated to calculate sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV) and diagnostic accuracy of NMP22, urine cytology and combined NMP 22 / urine cytology taking cystoscopy as gold standard. Receiver operating characteristic (ROC) curve analysis performed and the area under the curve (AUC) compared among these tests. Effect modifiers like age and gender were controlled by stratification. McNemar’s test used to compare sensitivity and specificity (P-value ≤0.05 was considered significant).

**RESULTS**

The study analyses included 380 patients. The average age of patients was 53.08± 12.41 years with age range (24-75) years and male to female ratio 3.75:1(300 males vs. 80 females). Rigid cystoscopy revealed BC in 215 (56.6%) patients while 165 (43.4%) were found negative. NMP 22 test showed BC in 207 (54.5%) patients and 173 (45.5%) had no cancer. Urine cytology was positive for TCC in 126 (33.2%) and negative in 254 (66.8%). In comparison to gold standard rigid cystoscopy, urine cytology revealed sensitivity 54% and specificity 93.9% (statistically significant p-value <0.001), NMP 22 had sensitivity 81.9% and specificity 81.2% (statistically significant p-value <0.001) while combination of both tests showed sensitivity 91.6% and specificity 77% (statistically significant p-value <0.001) (Table-I). Combination of NMP 22 / cytology outperformed NMP 22 and urine cytology in terms of sensitivity (91.63 vs 81.83 vs 53.96), NPV (87.59 vs 77.46 vs 61.02) and diagnostic accuracy (85.26 vs 81.58 vs 71.32) but at the cost of specificity (76.97 vs 81.21 vs 93.94) and PPV (83.83 vs 85.02 vs 92.06). Urine cytology revealed highest sensitivity (93.94%) and PPV (92.06%) while NMP 22 showed good balance of sensitivity, specificity, PPV, NPV and diagnostic accuracy (81.86% (76.05- 86.77), 81.21% (74.40-86.86), 85.02% (80.43- 88.69), 77.46% (71.93- 82.17), 81.58% (77.31- 85.35 respectively) (Table-II). Data was stratified according to age and sex for all three tests. Cytology performed equally in all age groups and both gender while NMP 22 had improved sensitivity and NPV in patients > 45 years of age. Combination of both performed better in female sub group (Table-III). ROC curve analysis revealed combination to have highest AUC (0.843 vs 0.815 vs .73) as compared to NMP 22 and Cytology, the difference being statistically significant (p-value <0.001) (Fig.1).

**DISCUSSION**

BC is highly common lethal disease affecting elderly posing very high financial and administrative burden on health care systems across the world. Rigid cystoscopies is criterion standard for diagnosis as well as surveillance nonetheless it is invasive, expensive and resource intensive. Urine cytology is another commonly used...
gold standard modality having excellent specificity but very low sensitivity especially for low grade tumors, requirement of trained cytopathologist and paucity of objective interpretation criteria. These compelling limitations prompted approval of several noninvasive biomarkers though with conflicting diagnostic accuracy quoted for primary and recurrent BC. The NMP-22 BladderChek test is an in vitro enzyme immunoassay approved by FDA for in-office testing of BC and harbors potentials to revolutionize BC diagnostic and surveillance strategies. In this prospective validational study we investigated the diagnostic value of urinary NMP 22 against urine cytology for BC.

Our data showed concordance with already published international data on the subject albeit with different study population. Stamfer et al. evaluated 231 patients with a history of CKD. They found a sensitivity of 64.4% and specificity of 84.8% for urine cytology. In our study, the sensitivity and specificity of urine cytology were 55.56% and 91.84%, respectively. The PPV, NPV, and accuracy of urine cytology were 57.91%, 51.11%, and 72.82%, respectively.

These results indicate that NMP-22 BladderChek test may have a potential role in the diagnosis of BC, and further studies are needed to confirm these findings.

Table III: Data stratification of sensitivity, specificity, PPV, NPV and accuracy of urine cytology, NMP-22 and Combined NMP-22 & Cytology with respect to age and gender (n=380).

| Test          | TP  | TN  | FP  | FN  | Sensitivity (95% CI) | Specificity (95% CI) | PPV (95% CI) | NPV (95% CI) | Accuracy (95% CI) |
|---------------|-----|-----|-----|-----|----------------------|----------------------|--------------|--------------|-------------------|
| Cytology Age  |     |     |     |     |                      |                      |              |              |                   |
| <45 Years     | 30  | 45  | 04  | 24  | 55.56% (41.4-69.08)   | 91.84% (80.4-97.73)  | 88.24% (74.9-95.18) | 65.22% (57.91-71.88) | 72.82% (63.16-81.12) |
| >45 Years     | 86  | 110 | 06  | 75  | 53.42% (45.4-61.3)   | 94.83% (89.08-98.08) | 93.48% (86.63-96.94) | 59.46% (55.29-63.5) | 70.76% (65.92-76.05) |
| Gender        |     |     |     |     |                      |                      |              |              |                   |
| Male          | 93  | 122 | 08  | 77  | 54.71% (46.9-62.34)  | 93.85% (88.23-97.31) | 92.08% (85.42-5.84) | 61.31% (57.18-65.28) | 71.67% (66.2-76.7) |
| Female        | 23  | 33  | 02  | 22  | 51.11% (35.77-66.3)  | 94.29% (80.84-99.3)  | 92% (74.39-97.85)   | 60% (52.39-67.15)   | 70% (58.72-79.74)  |
| NMP-22 Age    |     |     |     |     |                      |                      |              |              |                   |
| <45 Years     | 43  | 39  | 10  | 11  | 79.63% (66.47-89.37) | 79.59% (65.66-89.76) | 81.13% (70.88-88.37) | 78% (67.25-85.96)   | 79.61% (70.54-86.91) |
| >45 Years     | 133 | 95  | 21  | 28  | 82.61% (75.36-88.12) | 81.9% (73.67-88.43)  | 86.36% (81.04-90.37) | 77.24% (70.56-82.7) | 82.31% (77.3-86.62) |
| Gender        |     |     |     |     |                      |                      |              |              |                   |
| Male          | 139 | 106 | 24  | 31  | 81.76% (75.13-87.26) | 81.54% (73.79-87.8)  | 85.28% (80.03-89.33) | 77.37% (71.11-82.61) | 81.67% (76.82-85.88) |
| Female        | 37  | 28  | 07  | 08  | 82.22% (76.95 to 92.0) | 80% (63.06-91.56)   | 84.09% (72.88-91.22) | 77.8% (64.63-87.02)  | 81.25% (70.97-89.11) |
| Combined Age  |     |     |     |     |                      |                      |              |              |                   |
| <45 years     | 51  | 37  | 12  | 03  | 94.44% (84.6-98.84)  | 75.51% (61.13-86.66) | 80.95% (72.13-87.47) | 92.5% (80.24-97.4)  | 85.44% (77.12-91.6) |
| >45 years     | 146 | 90  | 26  | 15  | 90.68% (85.1-94.69)  | 77.59% (68.91-84.8)  | 84.88% (79.95-88.77) | 85.71% (78.58-90.75) | 85.2% (80.46-89.16) |
| Gender        |     |     |     |     |                      |                      |              |              |                   |
| Male          | 155 | 100 | 30  | 15  | 91.18% (85.86-94.98) | 76.92% (68.72-83.86) | 83.78% (79.7-87.65) | 86.96% (80.29-91.6) | 85% (80.45-88.84)  |
| Female        | 42  | 27  | 08  | 03  | 93.33% (81.73-98.6)  | 77.14% (59.86-89.58) | 84% (73.97-90.65)   | 90% (74.82-96.46)   | 86.25% (76.73-92.93) |
of superficial TCC of the bladder and found that NMP22 was two times more sensitive than cytology while Yafi FA et al.\textsuperscript{19}, Dogan C et al.\textsuperscript{20} demonstrated sensitivity of 70-80% for NMP22 test. In comparison, cytology showed sensitivity of ~10-40%. Our results revealed significantly higher sensitivity of NMP-22 than urine cytology (81.58 vs 53.96%; p value <0.001) while cytology was found more specific (81.83% vs 93.94%).

A study conducted by Pichler R et al.\textsuperscript{21} compared NMP-22 and urinary bladder cancer antigen (UBC) rapid to urinary cytology. Analysis of 75 patients showed sensitivity and specificity both NMP-22 and urinary cytology comparable (12.9% & 100% vs 25.8% & 100% respectively), the sensitivity both tests quite less as compared to our study while specificity comparable to our results. AUC for both was also quite less as compared to our data (0.63 cytology & 0.56 NMP-22 vs 0.73 cytology & 0.82 NMP-22).

In another study published BJUI in 2018, Pichler R et al.\textsuperscript{22} compared Xpert Bladder Cancer (BC) Monitor with cystoscopy and urinary cytology. The sensitivity, NPV and AUC of cytology was 0.33, 0.76 and 0.547. NPV value in our study for cytology was 0.6 while AUC was 0.730. The sensitivity, specificity and NPV of cytology depicted by Zhou L et al.\textsuperscript{23} are comparable to our results (30%, 91% & 86%). Another study conducted by Bier S. et al.\textsuperscript{24} included 758 bladders and 385 upper urinary tract (UUT) samples, the sensitivity of cytology & NMP 22 were found 74.6 & 100% for UUT and 59.3 & 62.5% for bladder samples while specificity was 66.6 & 50.9% for UUT and 82.9 & 31.3 for bladder samples.

However, the results published by Lotan Y. et al.\textsuperscript{25} contrary to our data. The sensitivity of NMP-22 BladderChek was 0.11 as compared to 0.82 while NPV was 0.86 (0.77 in our study). These differing results could be attributed to the different thresholds used in these studies and differences in the number of patients in each series. Our analytical results depicted superiority of NMP-22 to all other tests when all parameters of sensitivity, specificity, PPV, NPV and accuracy are taken into account together, a finding reported by Goodison S et al.\textsuperscript{26} as well.

Todenhofer T et al.\textsuperscript{27} studied diagnostic accuracy of combination of various markers with urine cytology. Their results showed very good performance for combination of cytology with FISH, uCyt+ but no additional affect by combining with NMP 22. However, the combination of both tests revealed a trade of significantly better sensitivity, NPV and accuracy at cost of low specificity and PPV in our study. To add further combination can act as very good screening tool owing to high sensitivity although in the wake of increasing cost and missing 9% of the cases. Although the combination of the NMP22 test and cytology cannot replace cystoscopy in the surveillance protocol at present, it can potentially reduce frequency of cystoscopic follow up reducing burden on patient as well as health care provider.

**Limitations of the study:** The results of present study should be interpreted with care as it’s a single center study, grade, stage as well as upper tract tumor were not taken into account leading to potential bias in interpretation of results. Moreover, cytopathologist involved were more than one which may have caused inter-observer variability.

**CONCLUSIONS**

NMP22 is a quick, point of care test having higher sensitivity, NPV and accuracy but similar specificity and PPV to urine cytology for detection of TCC urinary bladder. Combination outperformed both in terms of sensitivity while having modest specificity. At present these noninvasive markers can’t be recommended to replace cystoscopy however the NMP22 alone or in combination provides new avenues to adjunct cystoscopy in the diagnosis, screening and surveillance of BC.

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REFERENCES

1. Al-Husseini MJ, Kunbaz A, Saad AM, Santos JV, Salahia S, Iqbal M, et al. Trends in the incidence and mortality of transitional cell carcinoma of the bladder for the last four decades in the USA: a SEER-based analysis. BMC Cancer. 2019;19(1):46. doi: 10.1186/s12885-019-5267-3

2. Leal J, Luengo-Fernandez R, Sullivan R, Witjes JA. Economic burden of bladder cancer across the European Union. Eur Urol. 2016;69(3):438-447. doi: 10.1016/j.eururo.2015.10.024

3. Antoni S, Ferlay J, Soerjomataram I, Znaor A, Jemal A, Leal J, Luengo-Fernandez R, Sullivan R, Witjes JA. Economic burden of bladder cancer across the European Union. Eur Urol. 2017;71:96-108.

4. Kamat AM, Hanh NM, Efstatious JA, Lerner SP, Malmstrom PU, Choi W, et al. Bladder cancer. Lancet. 2016;388:2796-2810. doi: 10.1016/S0140-6736(16)30512-8

5. Scheller T, Hofmann R, Hegele A. Sex-related differences in urothelial cell carcinoma of the bladder in Germany. Cancer Manag Res. 2019;11:309-316.

6. Babajuk M, Bohle A, Burger M, Capoun O, Cohen D, Comperat EM, et al. EAU guidelines on non-muscle invasive urothelial carcinoma of the bladder: update 2016. Eur Urol. 2017;71:447-461.

7. Daneshmand S, Bazzargani ST, Bivalacqua TJ, Holzbeierlein JM, Willard B, Taylor JM, et al. Blue light cystoscopy for the diagnosis of bladder cancer: Results from the US prospective multicenter registry. Urol Oncol. 2018;36(8):361.e1-361.e6. doi: 10.1016/j.urolonc.2018.04.013

8. Gopalakrishna A, Longo TA, Fantony JJ, Owusu R, Foo WC, Dash R, et al. The diagnostic accuracy of urine-based tests for bladder cancer varies greatly by patient. BMC Urol. 2016;16:30.

9. Kassouf W. The value of urine cytology in the workup of hematuria. Cancer Cytopathol. 2016;124(3):303-304.

10. Chou R, Gore JL, Buckley D, Fu R, Gustafson K, Griffin JC, et al. Urinary Biomarkers for Diagnosis of Bladder Cancer: A Systematic Review and Meta-analysis. Ann Intern Med. 2015;163(12):922-931. doi: 10.7326/M15-0997

11. Badar F, Sattar A, Meerza F, Irfan N, Siddiqui N. Carcinoma of the urinary bladder in a tertiary care setting in a developing country. Asian Pac J Cancer Prev. 2009;10(3):449-452.

12. Human Kidney and Urine Proteome Project (HKUPP). Exosome preparation form urine. 2019. Available from: http://www.hkupp.org (Accessed 26th January 2020)

13. Alere Scarborough, Inc. 10 Southgate Road Scarborough, Maine 04074- Unites States T +1 (207) 730-5750. Available from: https://www.alere.com/en/home/product-details/nmp22-bladderchek.html {Accessed 26th January 2020}

14. Badar F, Sattar A, Meerza F, Irfan N, Siddiqui N. Carcinoma of the urinary bladder in a tertiary care setting in a developing country. Asian Pac J Cancer Prev. 2009;10(3):449-452.

15. Todenhofer T, Hennenlotter J, Esser M, Ding Y, Xu J, et al. Application of fluorescence in situ hybridization in the detection of bladder transitional-cell carcinoma: A multi-center clinical study based on Chinese population. Asian J Urol. 2019;6(1):114-121. doi: 10.1016/j.aajur.2018.06.001

16. Bier S, Hennenlotter J, Esser M, Mohrhardt S, Rausch S, Schwentner C, et al. Performance of Urinary Markers for Detection of Upper Tract Urothelial Carcinoma: Is Upper Tract Urine More Accurate than Urine from the Bladder? Dis Markers. 2018;2018:5823870. doi: 10.1155/2018/5823870

17. Lotan Y, O-Sullivan P, Raman JD, Shariat SF, Kavaleris L, Frampton C, et al. Clinical comparison of noninvasive urine tests for ruling out recurrent urothelial carcinoma. Urol Oncol. 2017;35:531.e15-531.e22.

18. Goodison S, Rossier CJ, Urquidi V. Bladder cancer detection and monitoring: assessment of urine- and blood-based marker tests. Mol Diagn Ther. 2013;17(2):71-84.

19. Yafi FA, Brimo F, Steinberg J, Aprikian AG, Tanguay S, Kassouf W. Prospective analysis of sensitivity and specificity of urinary cytology and other urinary biomarkers for bladder cancer. Urol Oncol. 2015;33(2):66.e25-31. doi: 10.1016/j.urolonc.2014.06.008

20. Dogan C, Petit ES, Yildirim A, Zemheri IE, Çanakçı C, Başok EK, et al. The value of the NMP22 test for superficial bladder cancer diagnosis and follow-up. Turkish J Urol 2013;39(3):137-142.

21. Pichler R, Tulchiner G, Fritz J, Schaefer G, Horninger W, Heidegger I. Urinary UBC Rapid and NMP22 Test for Bladder Cancer Surveillance in Comparison to Urinary Cytology: Results from a Prospective Single-Center Study. Int J Med Sci. 2017;14(9):811-819. doi: 10.7150/ijms.19929

22. Pichler R, Fritz J, Tulchiner G, Klinglmair G, Soleiman A, Horninger W, et al. Increased accuracy of a novel mRNA-based urine test for bladder cancer surveillance. BJU Int. 2018;121:29-37.

23. Zhou L, Yang K, Li X, Ding Y, Mu D, Li H, et al. Application of fluorescence in situ hybridization in the detection of bladder transitional-cell carcinoma: A systematic review and meta-analysis. Eur Urol. 2017;71:96-108.

24. Bier S, Hennenlotter J, Esser M, Mohrhardt S, Rausch S, Schwentner C, et al. Performance of Urinary Markers for Detection of Upper Tract Urothelial Carcinoma: Is Upper Tract Urine More Accurate than Urine from the Bladder? Dis Markers. 2018;2018:5823870. doi: 10.1155/2018/5823870

25. Lotan Y, O-Sullivan P, Raman JD, Shariat SF, Kavaleris L, Frampton C, et al. Clinical comparison of noninvasive urine tests for ruling out recurrent urothelial carcinoma. Urol Oncol. 2017;35:531.e15-531.e22.

Author’s Contribution:

MTS and MRZ conceived, designed and did statistical analysis & editing of manuscript. MTS, HU and SU did data collection. MRZ, HA, ZIM and KS reviewed, provided technical support and final approval of manuscript. All authors done critical revision of the manuscript for important intellectual content. All authors of this paper have equally contributed to this study, approved the final version to be published and are responsible for integrity of research.

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