ABSTRACT

Current regimens for the detection and surveillance of bladder cancer are invasive and have suboptimal sensitivity. Here, we present a novel high-throughput sequencing (HTS) method for detection of urine tumor DNA (utDNA) called utDNA CAPP-Seq (uCAPP-Seq) and apply it to 67 healthy adults and 118 patients with early-stage bladder cancer who had urine collected either prior to treatment or during surveillance. Using this targeted sequencing approach, we detected a median of 6 mutations per patient with bladder cancer and observed surprisingly frequent mutations of the PLEKHS1 promoter (46%), suggesting these mutations represent a useful biomarker for detection of bladder cancer. 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. In the surveillance setting, we detected utDNA in 91% of patients who ultimately recurred, with utDNA detection preceding clinical progression in 92% of cases. uCAPP-Seq outperformed a commonly used ancillary test (UroVysion, \(P = 0.02\)) and cytology and cystoscopy combined (\(P \leq 0.006\)), detecting 100% of bladder cancer cases detected by cytology and 82% that cytology missed. Our results indicate that uCAPP-Seq is a promising approach for early detection and surveillance of bladder cancer.

SIGNIFICANCE: This study shows that utDNA can be detected using HTS with high sensitivity and specificity in patients with early-stage bladder cancer and during post-treatment surveillance, significantly outperforming standard diagnostic modalities and facilitating noninvasive detection, genotyping, and monitoring.
INTRODUCTION

Bladder cancer is the sixth most common cancer in the United States, with an estimated 79,030 new cases in 2017 (1). After diagnosis and treatment for localized disease, the National Comprehensive Cancer Network guidelines recommend that patients undergo cystoscopy and urine cytology evaluation to monitor for recurrence every 3 to 6 months for 2 years and then at increasing intervals (1). Unfortunately, cystoscopy is invasive, and cytology has a low sensitivity, ranging from 20% to 53% (2). As a result of the need for this procedure-based, long-term follow-up, bladder cancer management costs more per patient lifetime than any other cancer (3).

Many attempts have been made to overcome these challenges by developing biomarkers for bladder cancer surveillance. The FDA has approved six different urine-based tests for bladder cancer recurrence, but they achieve modest sensitivities (55%–70%) and specificities (71%–83%), and none of the available tests have achieved widespread adoption (4).

Recent work has demonstrated the promise of analyzing circulating tumor DNA (ctDNA) in plasma to detect minimal residual disease (MRD) in a variety of tumor types (5–7). For example, recent work from our group has shown that hybrid capture–based ctDNA analysis using Cancer Personalized Profiling by Deep Sequencing (CAPP-Seq) can achieve 94% sensitivity and 100% specificity for detecting MRD post-treatment in plasma from patients with localized lung cancer (6, 8, 9). In the case of localized bladder cancer, analysis of tumor DNA in urine has been explored as a potential approach for detection and surveillance using amplicon-based high-throughput sequencing (HTS) approaches, with sensitivities and specificities ranging from 68% to 85% and 80% to 100%, respectively (10–13).

In this study, we developed a novel HTS-based hybrid-capture method for detection of urine tumor DNA (utDNA) called uCAPP-Seq and applied it to urine supernatant specimens. We hypothesized that assessment of utDNA would have superior performance characteristics compared with cytology for detecting early-stage bladder cancer and post-treatment residual disease.

RESULTS

Development of a Novel Assay for utDNA Detection

Given the practical challenges posed by purifying nucleic acids from large volumes of fluid, we adapted and optimized a previously described resin-based cell-free DNA (cfDNA) extraction protocol for urine samples (Fig. 1A), which performs as well as commercially available kits but allows analysis of larger volumes of urine (Supplementary Fig. S1A; ref. 14). We next established that in the presence of EDTA, urine cfDNA concentrations remain stable for at least 7 days at 4°C but not at room temperature (Supplementary Fig. S1B and S1C). Applying our optimized protocol to 185 urine samples with a median urine volume of 50 mL, we observed a median urine cfDNA concentration of 7.7 ng/mL and yield of 348.1 ng per sample (Supplementary Tables S1 and S2).

We next designed a custom capture panel for bladder cancer targeting recurrent single-nucleotide variants (SNV), insertions/deletions, and copy-number alterations. We began by including genomic regions covering known driver mutations in bladder cancer (15–17). We then applied our previously described algorithm to maximize patient coverage in the smallest possible genomic space using data from 412 bladder cancer cases from The Cancer Genome Atlas (TCGA; refs. 8, 18). The final panel covered ~311 kb of genomic space, included regions from 460 genes, and was predicted to identify a median of 7 mutations per patient with bladder cancer (ref. 18; Supplementary Table S3).

Urine contains a wider range of cfDNA fragment sizes than plasma (Supplementary Fig. S2A–S2C), necessitating modifications to the library preparation protocol and bioinformatic analyses that we had previously optimized for CAPP-Seq analysis of plasma. First, we tested if mutant DNA fragments from bladder cancers were enriched in short (<500 bp) or long (>500 bp) fragments of urine cfDNA. We observed similar variant allele fractions for driver mutations in patients with known bladder cancer in both size ranges (Supplementary Fig. S3A–S3C), indicating that size selection of DNA fragments prior to library preparation was unnecessary. Furthermore, we found that enzymatic fragmentation yielded significantly higher DNA recovery than acoustic shearing (Supplementary Fig. S3D; P = 0.0006) and increased the deduplicated sequencing depth over unfragmented urine cfDNA by approximately 2-fold (Supplementary Fig. S3E and S3F; P ≤ 2). Finally, we modified our previously published bioinformatics approach for analysis of plasma cfDNA to efficiently recover shorter DNA molecules present in fragmented urine cfDNA (see Methods).

To explore the utility of uCAPP-Seq, we applied our optimized protocol and the bladder cancer panel to 118 urine and 60 tumor samples from 130 patients with bladder cancer (Supplementary Tables S1, S2, and S4). These samples were derived from two independent patient groups (Methods; Supplementary Fig. S4), including one where urine was collected at the time of diagnosis (“Early Stage Bladder Cancer Group,” n = 54) and a second where urine was collected during surveillance after treatment for localized bladder cancer (“Surveillance Group,” n = 64).

In line with expectations from in silico predictions, we observed a median of 6 mutations per patient (Fig. 1B; Supplementary Tables S5 and S6). Next, we tested the concordance of mutations detected in tumor tissue and urine using 18 patients for whom paired urine and tumor tissue were available. Across these cases, a median of 66.7% of mutations that were found in tumor tissue were also identified in utDNA, whereas a median of 73.2% of mutations that were found in utDNA were also detected in paired tumor (Fig. 1C), with higher concordance between putative driver versus passenger mutations (P = 0.009). Within tumor tissue, mutations that were also found in urine had higher median allele fractions than those not found in urine (27.2% vs. 9.2%, P < 0.0001; Fig. 1D). Taken together, these results suggest that overall concordance between mutations found in bladder tumors and utDNA is high and is likely higher for truncal mutations than subclonal variants.

Tumor and Urine Genotypes in Bladder Cancer

Across the 81 patients for whom we profiled either tumor tissue or urine at time of bladder cancer diagnosis, the
The two most commonly mutated regions were the TERT and PLEKHS1 promoters (Fig. 2A). TERT promoter mutations were present in 74% of cases, with all mutations occurring at the two previously described hotspots (Fig. 2B; ref. 17). PLEKHS1 was originally described as a recurrent noncoding mutation in cancers by Weinhold and colleagues in an analysis of whole-genome sequencing data, with mutations at one of two single-nucleotide hotspots (hg19 chr10:g.115511590 and 115511593) found in 20 of 863 (∼2%) of cancers, including 8 of 20 of bladder cancers (17). We identified PLEKHS1 promoter mutations in 37 of 81 (46%) cases in our group, with all mutations clustered at these two previously described hotspots (Fig. 2B).

Nearly all other genes mutated in more than 10% of the bladder cancer tumors we profiled were well-characterized driver genes, including TP53, FGFR3, ERBB2, and RB1. Consistent with prior studies, we observed a significantly higher frequency of TP53 mutations in high-grade versus low-grade tumors (59.2% vs. 12.5%; P < 0.0001) and, conversely, found disproportionately more FGFR3 mutations in low-grade versus high-grade cases (59.4% vs. 10.2%; P < 0.0001; Fig. 2C; ref. 19). We also assessed copy-number variants across genes known to be significantly altered in bladder cancer that were adequately covered in our panel (16, 18) and found deletions involving chromosome 9p21 in 27% of cases, amplifications of ERBB2 in 7%, and deletions of RB1 in 2% (Fig. 2A; ref. 18).
Figure 2. Genetic findings across bladder cancers profiled in the study. A, Spectrum of genetic mutations and copy-number changes observed across 81 tumor and utDNA cases in this study, with clinicopathologic correlates. All tumor cases and all utDNA cases from patients with active cancer and at least one variant detected by genotyping were included in this analysis. All genes mutated in $\geq 10\%$ of cases are shown, as well as all genes evaluated for copy-number variants. The stage definitions follow the AJCC TNM Staging System for Bladder Cancer, 8th edition. B, Distribution of mutations in the TERT and PLEKHS1 promoters. C, Comparison of mutations across high-grade versus low-grade bladder cancers profiled in this study. D, Enrichment of the APOBEC mutational signature in the cfDNA of patients with active bladder cancer versus healthy controls. P values were calculated by multivariate regression controlling for total mutation count, median deduplicated sequencing depth, and the interaction between the two. APOBEC, apolipoprotein B mRNA editing enzyme, catalytic polypeptide-like.
We next tested the ability uCAPP-Seq to detect early-stage bladder cancer. We applied two different approaches to the detection of mutations in urine cfDNA, here designated “tumor-informed” and “tumor-naïve” profiling. The tumor-informed approach leverages prior knowledge by first sequencing a patient’s tumor specimen and germline tissue and then testing for the presence of these mutations in a urine sample using a Monte Carlo–based statistical approach, as previously described (8). In contrast, the tumor-naïve approach is designed to detect putative driver mutations without prior knowledge of the tumor genotype (see Supplementary Methods). We used 33 independent control subjects to establish threshold parameters for both the tumor-informed and tumor-naïve approaches, whereas 34 controls served to validate specificity (see Methods; Supplementary Fig. S6).

Our early-stage bladder cancer group consisted of pretreatment urine from 54 individuals with biopsy-proven bladder cancer (74% pTa, 6% pTis, 9% pT1, and 11% pT2) and 34 controls without bladder cancer. Although we attempted to match bladder cancer cases and non–bladder cancer controls for age and smoking status, controls were slightly younger (median 63.5 vs. 71) and contained fewer smokers (58% vs. 85%; Supplementary Tables S9 and S10). We observed a median urine cfDNA concentration of 12 ng/mL and yield of 523.3 ng among patients with active bladder cancer and a median urine cfDNA concentration of 6.88 ng/mL and yield of 254.5 ng among controls (P = ns for both comparisons).

Using the tumor-naïve approach, we detected 134 putative driver mutations in the urine of cases and only 1 in the controls (Fig. 3A). The 1 mutation detected in a patient who did not recur was a STAG2 nonsense mutation that was near the terminal end of the protein (residue 1029/1268) and therefore may not be a true driver. Cytology was positive in 37.8% of patients who developed recurrence, whereas urine cfDNA was positive in 84% (P = 0.0001; Fig. 4B and C). A concurrent multiplexed fluorescence in situ hybridization test for aneuploidy (UroVysion) was available for 7 patients who developed recurrent disease and was positive for 3 (42.9%). This sensitivity was comparable to cytology (P = ns) but significantly lower than tumor-naïve cfDNA profiling (P = 0.019; Fig. 4C). Among 32 cases where cystoscopy was performed at the same time as the utDNA specimen, gross tumor recurrence was diagnosed in 11 patients (34.4%), again resulting in significantly lower sensitivity than uCAPP-Seq (P < 0.0001). The sensitivity of cytology and cystoscopy combined (53.1%) was also significantly lower than tumor-naïve utDNA profiling (P = 0.0057). We detected mutant DNA in 1 of 27 patients who had at least 9 months of negative follow-up, resulting in a specificity of 96% (Fig. 4B). The specificity of cytology could not be assessed in the control group, as a positive cytology was an exclusion criterion.

Tumor-naïve profiling detected 100% of recurrent cases that had positive cytology (n = 14) and 73.9% that had negative cytology (n = 23; P = 0.04). We did not observe a significant difference in the detection rate between low-grade and high-grade tumors (Fig. 4D). In the subset of patients with available paired tumor tissue (n = 22), tumor-informed
Figure 3. Application of uCAPP-Seq to detect early-stage bladder cancer. A, Distribution of putative driver mutations identified in utDNA using tumor-naive profiling across patients with biopsy-proven bladder cancer (n = 54) and controls (n = 34), with associated tumor grade and cytology result. B, Receiver-operating characteristic analysis of tumor-informed profiling (n = 27 cases and 34 controls), tumor-naive profiling (n = 54 cases and 34 controls), and cytology (n = 50 cases and 18 controls). C, Correlates of detection by tumor-naive profiling among bladder cancer cases (n = 54). P values were calculated by the N-1 \( \chi^2 \) test for comparing proportions. D, Correlates of utDNA levels (haploid genome equivalents per mL, hGE/mL) among bladder cancer cases (n = 54). P values were calculated by the Mann–Whitney test. SNV, single-nucleotide variant; CNV, copy-number variant; Sn, sensitivity; Sp, specificity; AUC, area under the curve.
Figure 4. Application of uCAPP-Seq to detect residual disease in the surveillance setting. A, Distribution of putative driver mutations identified in urine cfDNA using tumor-naive profiling across cases that developed recurrent cancer (n = 37) and cases with at least 9 months of negative clinical follow-up (n = 27), with recurrent cancer defined by biopsy (32 cases) or alternative clinical evidence (5 cases), as specified in Supplementary Table S12. B, ROC analysis of tumor-naive profiling (n = 37 cases and 27 controls) and tumor-informed profiling (n = 11 cases and 11 controls) across surveillance group. C, Comparison of the sensitivities of cytology (n = 37), cystoscopy (n = 32), cytology plus cystoscopy (n = 32), UroVysion (n = 7), and tumor-naive profiling (n = 37) in detecting residual bladder cancer. D, Correlates of sensitivity for detecting disease by tumor-naive profiling. P values for E and D were calculated by the N-1 z^2 test for comparing proportions. E, Kaplan–Meier analysis of recurrence-free survival stratified by utDNA detection by tumor-naive and tumor-informed profiling (HR, 8.8 and 27.3), respectively and (F) by cytology (HR, 4.6). P values and HR were calculated by the log-rank test. G, Example of patient detected by tumor-naive profiling but missed by cystoscopy, cytology, and UroVysion, who was later diagnosed with muscle-invasive bladder cancer, requiring a radical cystectomy. SNV, single-nucleotide variant; CNV, copy-number variant; Sn, sensitivity; Sp, specificity; AUC, area under the curve; MRD, minimal residual disease.
Detection of Bladder Cancer by cfDNA Profiling

profiling achieved a 91% sensitivity (significantly improving on cytology, \( P = 0.002 \)) with 100% specificity (Fig. 4B). Tumor-naïve profiling, tumor-informed profiling, and cytology were all predictive of recurrence-free survival, but utDNA achieved a wider separation of outcomes in patients with positive and negative findings (HR, 8.8, 27.3, 4.6, respectively, all with \( P < 0.0001 \); Fig. 4E and F). No significant differences were observed between patients who recurred and those who did not for age, sex, smoking history, prior tumor stage or morphology, or prior treatment type (surgery vs. intravesical therapy). Patients who did not recur, however, were more likely to have had prior low-grade tumors and a longer interval between their last treatment and the specimen analyzed (Supplementary Tables S11 and S12). Nevertheless, in multivariate logistic regression including these parameters, tumor-naïve utDNA profiling results remained highly significant (\( P = 0.001 \)), with an adjusted odds ratio for a “positive” classification of 128 (Supplementary Table S13).

Detection of utDNA preceded clinical disease recurrence in 92% of patients by a median of 2.7 months. This lead time is likely an underestimate, because many patients who recurred in our cohort had earlier surveillance time points that were negative by cytology but that we did not have access to for utDNA analysis. In addition to allowing noninvasive detection, our approach may therefore also facilitate earlier diagnosis of recurrence. The potential utility of this approach is highlighted by a case that was clinically classified as negative by cytology, UroVysion, and cystoscopy at the landmark time point but as positive by tumor-naïve utDNA profiling (Fig. 4G). At the follow-up screening interval 6.1 months later, cytology was again negative but cystoscopy raised concern for recurrence. A transurethral resection of the bladder at 7.8 months revealed muscle-invasive bladder cancer, leading to a radical cystectomy at 11.7 months.

**DISCUSSION**

Here, we present a novel method for profiling utDNA called uCAPP-Seq and apply it to patients with bladder cancer. Our approach demonstrates high concordance for mutations between tumor and utDNA and enables genotyping of multiple somatic alteration types across a broad genomic space in a single integrated assay. Using this approach, we demonstrate that PLEKHS1 promoter mutations are among the most common somatic alterations in bladder cancer and are shed into the urine. We also highlight the potential of an APOBEC mutational signature as a utDNA biomarker for bladder cancer. In a group of patients with bladder cancer amenable to transurethral resection, we achieved a ~6-fold improvement in sensitivity over cytology while maintaining high specificity. Additionally, in a group of patients undergoing surveillance for recurrent bladder cancer after local therapy, we achieved ~2-fold improvements in sensitivity over cytology, the most commonly used ancillary test for bladder cancer (UroVysion), and gross evidence of disease by cystoscopy, detecting 92% of recurrences at a median of 2.7 months before clinical recurrence. Given the long lead time observed in some cases (Fig. 4G), it is tempting to speculate that intervention at the time of utDNA positivity might increase the likelihood of success of bladder-sparing interventions, though this will need to be tested in prospective trials. In summary, profiling of urine supernatants that are currently discarded in patients undergoing cytology could have significant value for disease detection and surveillance.

Across both groups, utDNA was detected in 21 of 21 (100%) bladder cancer cases that were positive by cytology and 54 of 66 (82%) bladder cancer cases that cytology missed. Although numerous studies have shown that ancillary molecular testing can provide added sensitivity when used in conjunction with urine cytology, currently available tests typically miss a substantial fraction of cases cytology detects and fail to achieve cytology’s high specificity (2, 4). Although uCAPP-Seq will need to be tested prospectively and in larger studies, our initial results suggest that it could offer a higher sensitivity alternative to cytology for the noninvasive detection of bladder cancer, for example, in the work-up of patients with microscopic hematuria. It could reduce the frequency of expensive and invasive cystoscopy procedures in a surveillance context. Although not explored in this study, uCAPP-Seq could potentially allow the monitoring of response to therapies such as intravesicular Bacillus Calmette-Guerin or neoadjuvant chemotherapy through frequent monitoring of utDNA levels. Additionally, our approach could enable repeated monitoring of genome evolution during treatment (20).

Several recent studies have explored the use of cellular or cell-free urine DNA as a potential biomarker for bladder cancer (10–13, 21). This includes two studies using amplicon-based HTS approaches (10, 12). Specifically, Ward and colleagues and Springer and colleagues evaluated DNA in the cellular fraction of urine samples, achieving 70% to 85% sensitivity and 93% to 97% specificity in identifying bladder cancer at diagnosis, and in the case of Springer and colleagues, 68% sensitivity and 80% specificity in identifying residual bladder cancer in a surveillance context (10, 12). Our approach differs in using hybrid-capture target enrichment, interrogating a significantly larger genomic territory and number of mutations per case, and detecting somatic variants and copy-number alterations in one assay. In contrast to Springer and colleagues, we analyzed urine cfDNA instead of cellular DNA. Focusing on urine cfDNA in the supernatant has the advantage of utilizing material that is currently discarded and that, in some cases, may contain higher variant allele fractions than DNA isolated from the cellular pellet (13, 21).

Limitations of our study include the case-control study design used for our surveillance group, which enriched for clinically lower-risk patients in the control subgroup compared with the cases. However, this imbalance has no impact on our comparison with cytology and UroVysion, as the same imbalances similarly affect those methods and would be expected to enhance their performance (2). Separately, the finding of utDNA positivity in the surveillance group remained highly significant in a multivariate analysis. Likewise, the cases and controls in our early-detection group were also slightly mismatched for median age and smoking status, raising the possibility that specificity might be lower in a perfectly matched control group. Finally, our cohort sizes are relatively small, and it will be important to test our method in larger patient cohorts and prospective trials.

In conclusion, we have developed a novel HTS-based approach to detecting utDNA and used it to explore genomic features and noninvasive diagnosis of bladder cancer. Our
approach substantially improves on the performance of cytology while maintaining high specificity. Importantly, our approach will need to be assessed in additional patient cohorts and prospective clinical trials to establish its clinical utility.

METHODS

Patient Selection and Sample Collection

Urine samples were collected from two groups, including (group 1) 54 patients with biopsy-proven early-stage bladder cancer and controls at the Veterans Affairs Palo Alto Healthcare System prior to cystoscopy. Samples were also collected from (group 2) 410 patients undergoing surveillance for recurrent bladder cancer between June 2016 and August 2017 from the Stanford Cytopathology lab, and from 33 healthy volunteers. Characteristics of samples and patients from each group are listed in Supplementary Tables S5 and S7, respectively. Across both groups, tumor and germline tissue was collected where available from formalin-fixed, paraffin-embedded tissue blocks. This study was conducted with Institutional Review Board approval from both institutions in accordance with the Declaration of Helsinki. Written informed consent was obtained for samples acquired within the Veterans Affairs Palo Alto Healthcare System. Informed consent was waived by the Institutional Review Board for the use of discarded samples acquired at Stanford Hospital.

Sample Processing

Samples were centrifuged to collect the pellet for clinical cytology, and the supernatant was combined with EDTA to a final concentration of 0.5 mmol/L. Samples were stored at 4 °C for up to 1 week, then at −80 °C. Among samples selected for further analysis, 10 to 50 mL of urine supernatant was combined with Q-Sepharose resin slurry after centrifugation for 3 minutes at 20,000 g. Residual urine manifold with 1 mL 2 mol/L LiCl in 70% ethanol, followed by 1 mL 0.5 mmol/L potassium acetate (pH 5.0) in 80% ethanol. The resin was then washed on Econo-Pac Chromatography Columns (Bio-Rad) with 10 mL of 0.3 mol/L LiCl/10 mmol/L sodium acetate (pH 5.5), and DNA was eluted with 1.675 mL of 2 mol/L LiCl/10 mmol/L sodium acetate (pH 5.5) into 5 mL 95% ethanol. The solution was then washed on a QIAquick column (Qiagen) on a vacuum manifold with 1 mL 2 mol/L LiCl in 70% ethanol, followed by 1 mL 75 mmol/L potassium acetate (pH 5.0) in 80% ethanol. Residual fluid was removed by centrifugation for 3 minutes at 20,000 × g prior to elution in Buffer EB (Qiagen). This protocol was adapted from Shekhtman and colleagues and scaled up for large fluid volumes (14). For each subject, a maximum of 42 ng of urine cfDNA (Supplementary Tables S5 and S7) was subjected to enzymatic fragmentation with the Kapa Hyperplus Kit (Roche) for 30 minutes at 37 °C, after which library preparation proceeded as previously described (8).

Panel Design and Hybrid Capture

Hybrid selection was performed with a custom SeqCap EZ Choice Library (Roche) designed through the NimbleDesign portal with support from the BioProd division, using genome build hg19/GRCh37. Hybrid capture and further processing were performed as described previously, prior to 151 × 2 bp paired-end sequencing on an Illumina HiSeq4000 (Illumina) with an 8 base indexing read.

Bioinformatics Pipeline

Raw reads were demultiplexed and subjected to quality control and adapter content removal with the AfterQC package (22), and then mapped to human genome assembly hg19/GRCh37 using BWA-MEM (https://arxiv.org/abs/1303.3997). Molecular barcoding, PCR duplicate removal, and adaptive variant calling were performed as previously described (8, 23) with modifications to support variable fragment lengths. Raw variant calls were subjected to removal of stereotyped technical artifacts calibrated on 12 healthy control urine cfDNA samples (i.e., “polished”), as previously described (9). APOBEC mutational signature enrichment was identified by the deconstructSigs R package.

Statistical Analyses

Sensitivity and specificity were assessed in ROC analyses, as described in the Supplementary Methods. The gold standard for a true positive was biopsy-proven cancer in all the cases in the early-detection cohort and in 52 of 37 cases in the surveillance cohort. There were 5 cases in the surveillance cohort that did not have a supporting biopsy but did have strong alternative evidence of recurrent disease, such as one or more “malignant” cytology diagnoses combined with a positive cystoscopy or imaging finding. Time-to-event analysis for recurrence-free survival was done using the log-rank test to estimate both P values and hazard ratios and expressed as Kaplan–Meier plots. All statistical analyses were done using Prism 7 (GraphPad Software), R v3.2.2 (http://www.r-project.org) through the RStudio environment, or medcalc.org. In calculating the sensitivity and specificity of cytology, “negative” and “atypical” results were considered clinically negative, whereas “suspicious” and “positive” results were considered clinically positive (23). See Supplementary Methods for further details.

Disclosure of Potential Conflicts of Interest

J.C. Dudley is a consultant/advisory board member for Merck. J.J. Chabon is a consultant/advisory board member for Lexent Bio. A.A. Chaudhuri is an advisor for Geneoxygen, LLC; has received commercial research support from Roche Sequencing Solutions; has received honoraria from the speakers bureaus of Roche Sequencing Solutions, Foundation Medicine, Inc., and Varian Medical Systems; and is a consultant/advisory board member for Geneoxygen, LLC, and Oscar Health. A.A. Alizadeh has ownership interest (including stock, patents, etc.) in CiberMed and FortySeven, and is consultant/advisory board member for Roche, Genentech, Chugai, Gilead, Janssen, Pharmacyclics, and Celgene. M. Diehn reports receiving a commercial research grant from Varian Medical Systems, has ownership interest in patents on cancer biomarkers, is a co-founder of CiberMed, and is a consultant/advisory board member for Roche and AstraZeneca. No potential conflicts of interest were disclosed by the other authors.

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Detection of Bladder Cancer by cfDNA Profiling

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