Molecular Characterization of Urothelial Carcinoma of the Bladder and Upper Urinary Tract

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Abstract

PURPOSE: A better understanding of the molecular basis of urothelial carcinoma (UC) is needed to refine the clinical decision-making process. METHODS AND MATERIALS: We performed next-generation sequencing to investigate the mutational and transcriptional profiles of commonly mutated genes in UC using Ampliseq v2. Copy number variations (CNVs) were detected with nCounter assay. Genetic alterations between upper tract UC (UTUC) and urinary bladder UC (UBUC) were compared. RESULTS: Tumor samples from 31 UTUC and 61 UBUC patients were included in analysis. The two groups showed similar clinicopathologic features including tumor grade and stage. Median survival was longer in UTUC than UBUC patients, though this was statistically nonsignificant (59 vs 41 months, \( P = .137 \)). In total, we found 982 genetic alterations from 92 samples: single nucleotide variants were the most common type of somatic mutation (479/508, 94.3%). Frequently detected somatic mutations included TP53 (68.5%), KDR (41.3%), and PIK3CA (17.4%). Notably, RB1 mutations were the only mutations significantly different between the UBUC and UTUC groups (19.7% vs. 0%, \( P = .020 \)). The most common types of CNVs included amplifications (56/62, 90.3%): 17.7% of patients identified amplifications in NOTCH1. We also identified five translocations in the entire study population, including one case with FGFR3-TACC3 (Chr4) fusion. CONCLUSION: Within a small study population, we identified similar genetic alterations in both UTUC and UBUC patients, indicating a basis for similar management strategies.

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Introduction

Urothelial carcinoma (UC), a cancer involving the transitional epithelium of the urinary tract, is the seventh most common malignancy in Korea [1]. The majority of UC arises in the urinary bladder (UBUC), whereas only 5% to 10% occurs in the upper urinary tract (UTUC) including the renal pelvis and ureter [2]. Because of the relative rarity of UTUC, clinical decision-making for patients with UTUC is based on treatment data for UBUC [3]. However, prognosis and treatment strategies vary between UTUC and UBUC. UTUC tends to have a
poor prognosis with a 5-year cancer-related survival <50% for pT2/pT3 tumors and <10% for pT4 tumors [4]. Although the treatment options for muscle-invasive UBUC have expanded in recent years to include (neo) adjuvant chemotherapy [5,6], no definitive recommendations exist regarding the use of perioperative chemotherapy in the management of UTUC. In both UTUC and UBUC, cisplatin-containing combination chemotherapy is considered standard treatment for patients with advanced metastatic disease.

There are few data supporting or refuting the assertion that we can apply similar principles to the management of UTUC and UBUC [7,8]. Although it is recognized that UTUC and UBUC harbor a similar morphology and cytogenetic changes as well as prognostic factors, controversy remains regarding whether UTUC and UBUC have similar biological behavior [9–11]. Molecular approaches are used extensively to enhance our understanding of cancer biology. The mutation landscape in muscle-invasive bladder cancer from The Cancer Genome Atlas (TCGA) suggests numerous therapeutic opportunities [12]. Gene expression signatures of muscle-invasive UBUC delineate tumor subtypes into luminal and basal types and are associated with efficacy to cisplatin-based chemotherapy [13]. Recently, immune therapy has shown considerable promise for the treatment of invasive UBUC [14]. Emerging immune biology data have revealed an association between response, immune check point inhibition, and survival and have established TCGA subtype and mutation load as potential biomarkers in UC [15,16]. Molecular studies have demonstrated some biological distinctions between UTUC and UBUC [17,18].

Understanding of the differences and similarities in the genetic landscapes of UTUC and UBUC is crucial to defining the utility of new diagnostic and treatment strategies. Based on these considerations, this study sought to characterize genetic alterations in UTUC and UBUC in Korean patients via next-generation sequencing (NGS) with Ampliseq.

Material and Methods

Patient Selection

This single-center, biomarker study is a part of the Samsung Medical Center (SMC) Oncology Biomarker study (ClinicalTrials.gov identifier: NCT01831609). We collected tumor samples from samples from 31 UTUC and 61 UBUC patients who were referred to our medical oncology department after radical cystectomy or neophrourectomy from 2012 to 2014. All patients provided written informed consent for the use of tumor tissues as well as their clinical data. This study was performed in accordance with the Declaration of Helsinki and approved by the Institutional Review Board of SMC (Seoul, Korea).

Genomic DNA Extraction

Our dedicated genitourinary pathologist (G.Y.K.) reviewed all pathology specimens to ensure the samples contained >80% tumor cells with <20% necrosis. Genomic DNA was extracted from formalin-fixed, paraffin-embedded (FFPE) samples using a QIAamp DNA Mini Kit (Qiagen, Valencia, CA). After extraction, genomic DNA quality and quantity were analyzed using a NanoDrop 8000 UV-Vis spectrometer (Thermo Scientific Inc., Willington, DE), Qubit 2.0 Fluorometer (Life Technologies Inc., Grand Island, NY), and 2200 TapeStation instrument (Agilent Technologies, Santa Clara, CA). Because this study was retrospective in nature, matched normal tissues were not available.

Results

Patients and Treatment Characteristics

Baseline characteristics of all patients are listed in Table 1. Median age at the time of surgery of all patients was 65 years (range, 37–83). UC patients were predominantly male (84.8%), but the proportion of female patients was significantly higher in UTUC than in UBUC (29.0% vs 8.2%, P = .02). According to SMC institutional guidelines, radical cystectomy and bilateral lymph node dissection (LND) is the standard treatment for patients with muscle-invasive or
locally advanced UBUC. For those with UTUC, unless enlarged lymph nodes are suspected before or during surgery, elective LND is not routinely performed. Therefore, all but one UBUC had undergone LND, whereas it was performed in only 48.4% of UTUC patients. There was significant difference in other clinico-pathological features including tumor grade, pT stage, and lymphovascular invasion between UBUC and UTUC.

Sixty-eight percent of UTUC patients received adjuvant chemotherapy after surgery. In the UBUC cohort, one-half of patients (49.2%) received neoadjuvant chemotherapy prior to radical cystectomy. In all patients, perioperative chemotherapy was a combination of gemcitabine plus either cisplatin or carboplatin, based on patient renal function.

Survival Outcomes

Median follow-up duration was 33 months (range, 28-37). Median overall survival was, although statistically insignificant, longer in the patients with UTUC compared with UBUC (59 months vs 41 months; $P = .137$; Figure 1). On multivariate analysis, only the presence of lymphovascular invasion was associated with decreased OS (hazard ratio, 3.34; 95% CI, 1.41-7.88; $P = .006$; Supplementary Figure 1).

Genomic Alteration Analysis Using the Ampliseq and Copy Number Variations

In total, we found 982 genetic alterations from 92 tumor samples (Supplementary Table 2). On average, there were 5.5 somatic mutations and 0.7 genomic copy numbers per sample. Figure 2 describes the landscape of genetic alterations for all patients with UC. Single nucleotide variants were the most common type of somatic mutation (479/508, 94.3%), followed by small insertion-deletions (indels; 29/508, 5.1%). On the other hand, the most common types of CNVs were amplifications (56/62, 90.3%) and deletions (6/62, 9.7%).

The frequency of mutations was not significantly different between UTUC and UBUC groups ($P = .13$). The median number of somatic mutations in UBUC and UTUC was 4 (range, 0-25) and 4 (range, 2-20), respectively. The most commonly observed mutation in UBUC patients was TP53 (67.2%) followed by KDR (44.3%), PIK3CA (19.7%), and RB1 (19.7%). In UTUC patients, TP53 (71.0%), KDR (35.5%), and TERT (16.1%) were the most frequently observed somatic mutations. Notably, we detected no RB1 mutations in the UTUC cohort compared with 12.9% frequency in UBUC tumors ($P = .020$). The most commonly observed CNV in UC patients was NOTCH1 (17.7%), followed by FGFR3 (14.5%) and MDM2 (12.9%). We found that the frequency of CNVs was not significantly different between the UTUC and UBUC groups. (See Fig. 3.)

We also identified five translocations in the total UC cohort including one case with FGFR3-TACC3 (Chr4) fusion that was already considered a promising therapeutic target. Other fusions with unknown biological significance included EWSR1-EMID1 (Chr22), ERBB2-PSMD11 (Chr17), ZNF507-AKT2 (Chr19) intrachromosomal translocations, and MLH1-FRY (chr3-chr13) interchromosomal translocations (Table 2).

Discussion

UTUC is a rare subset of UC with a poor prognosis that has not improved in recent decades, as the biological mechanisms of UTUC are still unclear. Whether we can apply similar principles in the treatment of UTUC based on UBUC also remains controversial. In the current study, we revealed that UBUC and UTUC shared common molecular features, while they also have site-specific features. Many observational
studies produced conflicting results regarding the significance of anatomical location on the prognosis of UC [19–21]. In the present study, nonsignificantly longer OS was observed in patients with UTUC compared with UBUC (59 months vs 41 months; \( P = .137 \)). Also, only the presence of lymphovascular invasion was associated with decreased OS on multivariate analysis (\( P = .006 \)).

However, it remains unclear whether the clinical behavior of UC originates from innate tumor biology. Recently, Blaveri et al. showed that the fraction genome altered was associated with worse outcome in muscle-invasive UBUC, independent of other clinicopathologic parameters [22]. Based on that study, we speculated that genetic aspects of the disease play an important role in the prognosis of patients with UC. Because of its rarity, comprehensive studies on the molecular basis of UTUC are scarce. Wu et al. identified ALDH2, CCNE1, and SMAD3 as potential prognostic markers in UTUC [23]. Sanford et al. found that UTUC tended to exhibit high expression of genes such as SLITRK6 associated with a luminal subtype [24]. In the current study, the landscape of alterations in UTUC was similar to that of UBUC. Consistent with the genetic alterations by TCGA and others, specific genes including TP53, PIK3CA, and FGFR3 were the main molecular alterations associated with both UBUC and UTUC [12,17,18]. TP53 was the most frequently mutated gene (68.5% of all UC cases). On the other hand, FGFR3 mutations (13.0%) occurred less frequently. This variation in genetic alterations can be explained by the higher prevalence of high-grade tumors in this study population. Bakkar et al. demonstrated that FGFR3 mutations were associated with low-stage, low-grade tumors, whereas TP53 mutations were associated with high-stage, high-grade tumors [25]. Surprisingly, KDR mutations were detected in 35.5% of UTUC and 44.3% of UBUC cases in our study. The KDR gene encoding for VEGFR-2 is considered a significant prognostic marker in colorectal carcinoma [26]. However, no prior studies have evaluated the role of KDR mutations in UC. Millis et al. showed that only 2% of UBUC patients harbored KDR mutations [27]. Further study with NGS to analyze both somatic and germline variants in KDR genes is needed. It is noteworthy that the prevalence of mutations differed. The Rb1 gene was the only mutated gene in UBUC. Interestingly, Rb1 mutations were significantly associated with UBUC. Similarly, comparison of UTUC with UBUC by Sfakianos et al. revealed a lower prevalence of mutations in Rb1 [17].

In this study, the most common amplifications were NOTCH1 (17.7%) and FGFR3 (14.5%). The molecular basis for the NOTCH1
amplifications in UC needs to be further elucidated. While FGFR3 has long been considered an attractive therapeutic target in UC, little is known about the role of FGFR1. Tomlinson et al. demonstrated that FGFR1 expression is increased in UBUC tissues and promotes cell proliferation and survival via activation of the MAPK pathway in UBUC cell lines [28]. In the current study, FGFR1 amplification was only observed in the UTUC group (3.2%). Further work will be needed to validate the relative significance of different candidates according to the tumor location and to evaluate their interplay.

Although muscle-invasive UBUCs show many chromosomal rearrangements, the only recurrent gene-gene fusion that has been identified is FGFR3 gene rearrangement [29]; FGFR3 with one of two different fusion partners: TACC3 or BA1AP2L1 (BA1-associated protein 2-like 1). Both FGFR3-TACC3 and FGFR3-BA1AP2L1 translocations generate constitutively activated and oncogenic FGFR3 kinase protein products, and cellular dependence on these drivers confers sensitivity to selective FGFR inhibitor [29,30]. In this study, we identified one case of FGFR3-TACC3 fusion. In addition, we noted the presence of several novel genomic rearrangements including EWSR1-EMID1, ERBB2-PSMD11, SNF507-AKT2, and MLH1-FRY. However, whether this genomic rearrangement produces an in-frame fusion event and the functional significance of these novel rearrangements have yet to be established.

The present study has several limitations. First, the results should be interpreted with caution given the retrospective nature and limited number of samples and mutations analyzed in the current study. Second, accurate identification of somatic mutations was challenged in the absence of normal DNA to distinguish somatic mutations from germline polymorphisms. Third, epigenetic differences and/or differences in gene expression may be more important drivers of disease progression than genomic alterations. Lastly, to extract biologically relevant information from molecular alterations, functional validation is needed.

In conclusion, UC is biologically heterogeneous and has widely variable clinical outcomes and responses to conventional chemotherapy. Our study characterized similarities and differences in the patterns of genetic alteration between UBUC and UTUC. A comprehensive understanding of the biology of UTUC is needed to identify new drug targets in order to improve clinical outcomes.

Supplementary data to this article can be found online at https://doi.org/10.1016/j.tranon.2017.10.008.

Disclosure of Potential Conflicts of Interest

There are no conflicts of interest to declare.

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