Whole-exome sequencing identified mutational profiles of urothelial carcinoma post kidney transplantation

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Abstract
Kidney transplantation is a lifesaving option for patients with end-stage kidney disease. In Taiwan, urothelial carcinoma (UC) is the most common de novo cancer after kidney transplantation (KT). UC has a greater degree of molecular heterogeneity than do other solid tumors. Few studies have explored genomic alterations in UC after KT. We performed whole-exome sequencing to compare the genetic alterations in UC developed after kidney transplantation (UCKT) and in UC in patients on hemodialysis (UCHD). After mapping and variant calling, 18,733 and 11,093 variants were identified in patients with UCKT and UCHD, respectively. We excluded known single-nucleotide polymorphisms (SNPs) and retained genes that were annotated in the Catalogue of Somatic Mutations in Cancer (COSMIC), in the Integrative Onco Genomic cancer mutations browser (IntOGen), and in the Cancer Genome Atlas (TCGA) database of genes associated with bladder cancer. A total of 14 UCKT-specific genes with SNPs identified in more than two patients were included in further analyses. The single-base substitution (SBS) profile and signatures showed a relative high T > A pattern compared to COMSIC UC mutations. Ingenuity pathway analysis was used to explore the connections among these genes. GNAQ, IKZF1, and NTRK3 were identified as potentially involved in the signaling network of UCKT. The genetic analysis of posttransplant malignancies may elucidate a fundamental aspect of the molecular pathogenesis of UCKT.

Keywords: Kidney transplantation, Malignancy, Urothelial carcinoma, Whole exome sequencing, Mutations

Introduction
Kidney transplantation is a lifesaving option for patients with end-stage kidney disease (ESKD) because it significantly improves survival; it has a mortality rate that is 63–80% lower than that of continued dialysis [1, 2]. Modern immunosuppressive agents have tremendously reduced the incidence of acute rejection within the first year and achieved outstanding short-term patient and graft survival [3]. However, the long-term survival rate of patients who receive kidney transplants is low. Recipients of a kidney transplant experience various complications, including cardiovascular disease and posttransplant malignancies (PTMs), that lead to morbidity and mortality [3].

The incidence of cancer in recipients of a kidney transplant is at least 2 to 4 times higher than that of their age-matched and gender-matched counterparts in the general population [4–6]. Studies from Western countries have reported that nonmelanoma skin cancer is the most common PTM [7]. The incidence of skin cancer is lower in Asian countries. Urothelial
carcinoma (UC) of the bladder and urinary tract is the most common PTM in Taiwan and Mainland China, comprising approximately 40% of PTMs [8, 9]. In a nationwide cohort population-based study, Tsai et al. reported that recipients of a heart, lung, kidney, or liver transplant exhibited a risk of de novo cancer that was three times higher than that of the general population [10]. The standardized incidence ratio of urinary tract malignancies (among which bladder cancer was the most common) among recipients of a kidney transplant was 10.93 (95% CI, 9.20–12.99) [10]. Furthermore, in our previous single-center retrospective study, the most common PTMs were UC and hepatocellular carcinoma [11].

UC has a high mutational burden and exhibits greater molecular heterogeneity than do other solid tumors [12]. Different genetic alterations and pathogenic pathways occur in UCs at various anatomical locations [13, 14]. Somatic genetic mutation is a highly influential factor in UC tumorigenesis and progression. An abundance of information regarding somatic alteration in UC has been published due to rapid progress in next-generation sequencing [15]. According to the Cancer Genome Atlas (TCGA) database, the genes with the most common mutations in bladder UC are TP53, KDM6A, and ARID1A [15, 16]. TP53 mutations were commonly observed in high-grade tumors, whereas FGFR3, CREBBP, and STAG2 mutations were more commonly observed in low-grade tumors [15].

Recipients of a kidney transplant are prone to developing cancer due to underlying diseases leading to renal failure, chronic infections by oncogenic viruses, immunocompromised status related to treatment with immunosuppressive drugs, nutritional deficiencies, or altered deoxyribonucleic acid (DNA) repair. Knowledge of the underlying somatic genomic alterations in UC developed after kidney transplantation (UCKT) is limited. To our knowledge, no unbiased systematic effort has been made to describe genomic alterations in UCKT.

Whole-exome sequencing (WES) has been extensively applied to identify drivers and somatic alterations in diseases, including cancer genome profiles, in both research and clinical settings [17, 18]. WES is an effective tool for medical genetic research because it targets almost all protein-coding regions in the human genome. In this study, we performed WES and analyzed UC tissues from recipients of a kidney transplant (the UCKT group) and patients on hemodialysis (the UCHD group) to further characterize the unique genomic landscape of UCKT.

Materials and methods

Sample collection and DNA extraction

The sample collection and study protocol were conducted under the Institutional Review Board of Kaohsiung Medical University Hospital (KMUHIRB-G(I)-20150030). Written informed consent was obtained from all the patients, and all clinical investigations were conducted in accordance with the Declaration of Helsinki.

A total of 12 tumor samples, comprising 7 formalin-fixed archival UC samples from 6 recipients of a kidney transplant and 5 UC samples from patients on hemodialysis, were collected. Recipients of a kidney transplant who had a diagnosis of UC before their respective transplants were excluded. Because chronic kidney disease (CKD) and ESKD are risk factors for UC, five patients on hemodialysis who received a diagnosis of UC after the commencement of dialysis were selected to form the control group. Each patient’s medical history was confirmed by chart review. None of the patients had previously received chemotherapy or radiotherapy.

After a pathologist reviewed all the samples, the tumor samples from the paraffin blocks were macrodissected. All the specimens were placed in Eppendorf containers to prevent tissue cross-contamination. Genomic DNA was extracted using a FavorPrep formalin-fixed paraffin-embedded (FFPE) Tissue DNA Extraction Micro Kit (Favorgen Biotech, Pingtung, Taiwan) from FFPE tumor tissue. The DNA was quantified and qualified using Qubit 4 Fluorometer (Thermo Fisher Scientific, Waltham, USA), 4150 TapeStation System (Agilent, Santa Clara, USA), and Nanodrop spectrophotometer (ASP-2680, Celltagen, Seoul, Republic of Korea) according to the manufacturer’s protocol. One microgram DNA from each tumor sample was subjected to run the WES analysis.

Whole-exome sequencing

Library enrichment for WES was conducted using a SureSelectXT V6_r2 reagent kit (Agilent Technologies, Santa Clara, CA, USA). The enriched samples were sequenced using an Illumina HiSeq2500 system (San Diego, CA, USA) with a 2 × 76–base pair (bp) paired-end sequencing approach. The mean coverage of the exomes was 79.76 × , and more than 96% of the exomes had a coverage higher than10 × . The sequencing data were annotated according to the GRCh37/hg19 reference genome.

The data were examined using a data analysis pipeline developed at the Cologne Center for Genomics (CCG).
[19]. Initially, raw sequencing reads were mapped to human genome reference (hg19) using Burrows-Wheeler Alignment tool (BWA) [20] followed by duplication marking using Picard [21].

Subsequently, base quality score recalibration and local indel realignment were performed according to Genomic Analysis Toolkit (GATK) practice guideline [22, 23]. After the post-alignment improvements, variant calling was performed using GATK’s unified-genotyper and SAMtools’ mpileup [24]. The variant lists from both callers were merged and annotated using various public databases: dbSNP [25], 1000 Genomes Project [26], Exome Variant Server [27], HGDM professional database [28], dbVAR and DGVa [29], GERP, and Ensembl [30]. The functional effect of variants were predicted using PolyPhen [31], SIFT [32], and RVIS [33]. All of these downstream analysis has been assembled together with a few scripts developed in-house. Splice site analysis based on Yeo [34] et al. was performed. Finally, the annotated variant list is uploaded to CCG’s web interface and VARBANK database [35]. Scripts developed in-house at the Cologne Center for Genomics were applied to detect protein changes, affected donor and acceptor splice sites, and overlaps with known variants. Our analysis focused on single-nucleotide variants/polymorphisms (SNVs or SNPs) and insertions or deletions (InDels) that may result in alterations in primary protein structure or in strong splice site effects [34].

Identifying specific genes in UCKT

The processed results of all 12 samples were then downloaded from VARBANK analysis platform. A series of steps were performed to identify cancer driver genes that contain novel mutations and are not reported previously in UCKT patients. First, mutations with dbSNPs annotations and with low allele frequencies (≤ 5%) were excluded from the data sets.Mutations at the same genomic location that were annotated as different transcripts were counted as one occurrence.

To ensure that these genetic alterations identified in our study were associated with tumor development, we identified genes with these mutations and using these gene names to search against the Catalogue of Somatic Mutations in Cancer (COSMIC) [36]. If in our list, a gene is also a reported cancer driver gene in COSMIC, the gene and its mutations was retained for further analysis. In other words, these newly identified mutations in our samples were considered potential driver mutations.

Subsequently, we utilized previous studies or databases that had analyzed bladder cancer including the COSMIC [36], the Integrative Onco Genomic (IntOGen) cancer mutations browser [37], the TCGA bladder tumor database [15], and the Chinese bladder cancer genome [38] to compare with our data. Cancer driver genes that were retained in the previous step but had not been reported in these bladder cancer studies were of great interest.

Then we selected the above driver genes which were identified in UCKT but not in UCHD groups. Finally, we reported genes that were found in two or more UCKT
patients. Figure 1 illustrates the data processing steps and corresponding mutations and gene numbers for UCKT and UCHD groups. COSMIC UC single-base mutation data was retrieved from the Cancer Browser in COSMIC [39] by using the selection of Primary site (Urinary tract), Sub site (Bladder), Histology (Carcinoma) and Sub histology (Transitional cell carcinoma). The mutational signatures were conducted using the SigProfiler tools (MatrixGenerator, Extractor and Plotting) [40]. Data from UCKT, UCHD and COSMIC UC were applied separately.

Sanger sequencing analysis for confirmation of somatic mutations
Genomic DNA from patients’ peripheral blood mononuclear cells (PBMC) was used as reference for detecting somatic mutations. We used a FavorPrep Blood / Cultured Cell Genomic DNA Extraction Mini Kit (Favorgen Biotech, Pingtung, Taiwan) to extract the genomic DNA from PBMC. All the somatic mutations of the selected genes were independently verified through Sanger sequencing of the PBMC genomic DNA. Primer pairs were designed using the University of California, Santa Cruz Genome Browser [41] and Primer-BLAST Browser of the National Institutes of Health [42].

The target regions were investigated using a touchdown polymerase chain reaction (PCR) protocol. The PCR mixture contained primers at 5 μmol L⁻¹ each, Taq DNA polymerase (Vazyme Biotech, China), and 50 ng/ul genomic DNA. The following touchdown PCR protocol was executed in a T100 thermal cycler (Bio-Rad Laboratories, CA, USA): initial denaturation at 94 °C for 5 min; annealing temperature decreased by 0.5 °C every other cycle: 94 °C for 30 s, annealing from 72 to 55 °C for 30 s, 72 °C for 30 s; followed by a cycle of 94 °C for 30 s, 55 °C for 30 s, and 72 °C for 30 s; and extension at 72 °C for 10 min. The PCR products were resolved in 1.5% (w/v) agarose gel for 20 min at 100 V in 1 × TAE buffer and stained with Gel Red biosafety dye (Biotium, CA, USA). The gel was documented on a UV transilluminator (Major Science, Taiwan). The DNA sequencing was performed at the core facility of the National Yang Ming Chiao Tung University (NYCU) Genome Center.

Analysis of gene expression profile by ingenuity pathway analysis
The Ingenuity Pathway Analysis (IPA version 68752261; QIAGEN) tool was used to establish protein networks. The main molecules detected from WES analysis and Sanger sequencing verified from the PBMC genomic DNA PCR products were set as the focusing molecules and were analyzed using the build tool to determine how their relationships were affected by UC. The molecules from the QIAGEN knowledge base were then added to the network according to their specific ontological associations.

Copy number variation (CNV) detection
CNV analysis was performed using ExomeDepth (ver 1.1.15) R software package, which uses read depth data to call CNVs from exome sequencing experiments. We generated read count data from WES BAM files, and compared the count results with an aggregate reference made up of samples from the same sequencing run to determine copy number at exon level resolution. Data were analyzed according to the ExomeDepth standard procedure [43].

Results
Baseline characteristics
The baseline characteristics of all the patients and their clinical and pathological features are listed in Table 1. All patients were female and sporadic, with no family history of UC nor any history of smoking. The majority (10 out of 12) had high-grade tumors. The KT80 and KT721 samples were from the same patient, a woman who underwent surgical removal of different parts of her genitourinary tract system over 5 years (Table 1). In the KT recipients, all the tumors occurred in their native genitourinary systems. The patients underwent induction treatments with steroids and IL-2 receptor antagonists. Tacrolimus, mycophenolate mofetil, and steroids were selected as maintenance agents. The average age at UC diagnosis was younger in the UCKT group than in the UCHD group (55 vs. 61.6 years, respectively; Table 2).

Whole-exome sequencing result analysis of UC
After WES was performed, the results were aligned to the human reference genome (hg19) and further used in a bioinformatics analysis. At least 100 million reads were collected from each WES sample. In the UCKT and UCHD groups, the mean coverages were 81 and 77 reads for UCKT and UCHD, respectively, and 96.18% and 95.78% of the coding exons, respectively, were covered by at least 10 reads (Table 2). A total of 18,733 and 11,093 variants were identified from the UCKT and UCHD cohorts (Fig. 1).

The majority of the mutations were unique to individual patients. Mutations were slightly more common in the UCKT group than in the UCHD group. An analysis of the WES data revealed an average of 564 known SNPs per individual. To identify new UC-related alterations, the mutations with dbSNP annotations were excluded.

We used the COSMIC cancer driver genes database to identify new SNPs in our cohorts. During this step, 278 and 235 genes in the UCKT and UCHD groups,
### Table 1 Baseline characteristics and clinic-pathologic features of all patients

| Patient ID | Gender | Age at diagnosis | HBV/HCV | Chinese Herb history | Occupation | Kidney transplant | Induction | Maintenance | Acute rejection | Location | Tumor pathology | Treatment |
|------------|--------|------------------|---------|----------------------|------------|-------------------|-----------|--------------|----------------|----------|----------------|-----------|
| KT17       | F      | 48               | No      | Unknown              | Bio-engineer | Yes               | Tac       | MMF + Pred   | Yes            | Bladder  | Infiltrating UC, high grade, pT3NoMx | Partial cystectomy |
| KT79       | F      | 61               | No      | Yes                  | Housewife   | Yes               | Tac       | MMF + Pred   | Yes            | Bladder, ureter | Non-invasive UC, high grade, pTa | Left nephrectomy |
| #KT80      | F      | 51               | No      | Yes                  | Housewife   | Yes               | Tac + MMF | + Pred + MTOR | No             | Ureter (l.t) | Infiltrating UC, high grade, pT2N0Mx | Bilateral nephrectomy |
| KT81       | F      | 63               | HBV     | Unknown              | Housewife   | Yes               | Tac       | MMF + Pred + MTOR | No             | Renal Pelvis | Infiltrating UC, high grade, pT3 | Left nephrectomy |
| KT720      | F      | 46               | No      | Unknown              | Medical      | Yes               | Tac + MMF | + Pred + MTOR | No             | Renal Pelvis | Non-invasive papillary UC, high grade, pTa | TURBT |
| #KT721     | F      | 55               | No      | Unknown              | Housewife   | Yes               | Tac + MMF | + Pred + MTOR | No             | Bladder | Infiltrating UC, high grade, pT2aN0Mx | Left nephrectomy |
| KT722      | F      | 61               | No      | Unknown              | Housewife   | Yes               | Tac + MMF | + Pred       | No             | Renal Pelvis | Infiltrating UC, high grade, pT2aN0Mx | Left nephrectomy |
| KT73       | F      | 52               | No      | Unknown              | Housewife   |                   |           |              |                | Bladder, ureter | Non-invasive papillary UC, low grade | Right nephrectomy and cystectomy |
| KT77       | F      | 56               | HCV     | Yes                  | Housewife   |                   |           |              |                | Bladder | Infiltrating UC, high grade, pT1Nx | Radical cystectomy and bilateral nephrectomy |
| KT78       | F      | 70               | No      | Unknown              | Sales person | No               | N/A       |              | N/A            | Bladder + ureter | Infiltrating UC, high grade, pT2aN0Mx | Radical cystectomy |
| KT595      | F      | 64               | No      | Yes                  | Worker      |                   |           |              |                | Renal Pelvis | Infiltrating UC, low grade, pT1NxMx | Left nephrectomy |
| KT596      | F      | 66               | No      | Unknown              | Housewife   |                   |           |              |                | Bladder | Non-invasive papillary UC, high grade | Left nephrectomy |

Abbreviations: UC, Urothelial carcinoma; Tac, Tacrolimus; MMF, Mycophenolate myfortil; Pred, prednisolone; MTORi, Mammalian target of rapamycin inhibitors. N/A: not applicable

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#KT80 and KT 721 sample are from the same patient at different time of surgery
respectively, were retained (Fig. 1). To identify onco-
driver genes specific to our cohorts (i.e., previously
unknown to be related to UC), we first collected UC-
related studies from the COSMIC [36], IntOGen cancer
mutations browser [37], TCGA bladder tumor database
[15], and Chinese bladder cancer genome [38] and we
excluded genes that had already been reported in these
studies. Ultimately, we identified 150 and 121 genes in
the UCKT and UCHD groups, respectively, that had
never previously been known to be associated with UC.
Of these cancer driver genes, 79 were uniquely identi-
fied in the UCKT cohort, and 17 of them (BTK, CARD11,
ELL, FNBPI, GNAQ, HOXD13, IKZF1, MAX, MLLT10,
NTRK3, PAX5, SEPTIN6, SEPTIN9, SH3GL1, SLC34A2,
TAL1, and TRAF7) exhibited mutations that occurred in
two or more patients (Table 3).

To verify the tumorigenic nature of the 14 genes in
the UCKT cohort (CARD11, FNBPI, GNAQ, HOXD13,
IKZF1, MAX, MLLT10, NTRK3, SEPTIN6, SEPTIN9,
SH3GL1, SLC34A2, TAL1, and TRAF7), we developed
a reliable PCR system to validate the mutations in the
patients’ PBMC genomic DNA and subjected to Sanger
sequencing analysis. The results revealed that mutations
in GNAQ, SEPTIN6, NTRK3, and IKZF1 occur only in
tumor samples. The possible interactions among these
genes deserved further pathway analysis.

SBS and CNV analysis
The mutations were classified into two categories: single-
base substitutions (SBSs) and InDels (insertions/dele-
tions) (Table 2). The single-base substitution category
was further divided into six subcategories: synonymous,
missense, nonsense, frameshift, splice acceptor or donor,
and nonstop (Table 2). The frameshift subcategory com-
prising one-base-pair differences between the alleles and
reference sites, whereas the InDels category comprises
mutations involving longer regions. Missense muta-
tions were the most common type in both the UCKT
and UCHD groups (74.75% and 79.40%, respectively;
Table 2). These mutations are associated with amino
acids changes in protein sequences, thereby affecting
the structures and functions of the proteins that their

Table 2 Summary of whole-exome sequence analysis of human urothelial carcinomas in kidney transplantation (UCKT) and
hemodialysis (UCHD)

|                                | UCKT (N = 7) | UCHD (N = 5) |
|--------------------------------|-------------|-------------|
| Average age at diagnosis (years) | 55          | 61.6        |
| Mean coverage                  | 81          | 77          |
| Targeted bases with at least 10 reads (%) | 96.19 | 95.78 |
| Known SNPs identified in targeted region* | 4198 | 2574 |
| Total somatic mutations *      | 18,703      | 11,060      |
| Insertion or deletion          | 271         | 188         |
| Single based substitution      | 18,432      | 10,872      |
| Synonymous                     | 1976        | 65          |
| Missense                       | 13,980      | 8782        |
| Nonsense                       | 948         | 550         |
| Frameshift                     | 288         | 186         |
| Splice acceptor                | 904         | 923         |
| Splice donor                   | 301         | 331         |
| Nonstop                        | 35          | 35          |
| C > A                          | 7.03        | 6.72        |
| C > G                          | 7.74        | 6.78        |
| C > T                          | 27.69       | 23.96       |
| T > A                          | 40.03       | 44.63       |
| T > C                          | 11.80       | 12.06       |
| T > G                          | 5.70        | 5.86        |

The WES results of the 12 samples were aligned to the human reference genome (hg19) and further subjected for bioinformatics analysis. UCKT; N = 7 and UCHD; N = 5
*The number was calculated as the summation of the samples in each group
**The number was calculated as the average of the samples in each group
respective genes encode. Interestingly, synonymous substitutions were much more common among the patients with UCKT than among those with UCHD (10.57% vs. 0.59%), which might indicate that the overall mutation rate increases after kidney transplantation, resulting in a higher risk of cancer. Among the nucleotide changes, T > A was the most common one in both the UCKT and UCHD groups (Table 2). In addition, in a TCGA study, 51% of all the mutations in bladder cancers were TpC > T or G mutations and 3.8% were TpC > A mutations [15]. Figure 2 illustrates the mutational features of the patients. Although missense mutations were the most common somatic mutations, followed by nonsense mutations, splice site mutations were also common. The effects of splice site mutations are difficult to determine; however, they may affect protein function by causing alternative splicing, intron retention, or exon skipping.

To further analyze the mutation signatures and compare to other cohort, total of 95,657 unique mutations were obtained from the Cancer browser in COSMIC for UC. The distribution of six classes of SBS showed distinct patterns among UCKT, UCHD and COMIC UC data (Fig. 3). The UCKT and UCHD showed relative high percentage in T > A (40.03% and 44.63%) while COSMIC showed high C > T (42.12%) and C > G (35.17%) mutations.

The results from SigProfiler tool [44] showed that both SBS5 in UCKT and UCHD presented highly similar to the SBS22 signature (83.6% and 81.12%) with the proposed etiology “Aristolochic acid exposure”, SBS5 (10.44% and 11.18%) with “unknown aetiology mutational burden is increased in bladder cancer samples with ERCC2 mutations and in many cancer types due to tobacco smoking” and SBS1 (5.96% and 7.7%) with “an endogenous mutational process initiated by spontaneous or enzymatic deamination” while COSMIC UC is similar to SBS5 (27.12%) SBS1 (4.48%), SBS13 (40.36%) and SBS2 (28.04%), respectively (Additional file 1: Fig. S1). These findings might indicate the possible unique etiology for the UC development in Taiwan.

In addition, CNV data was retrieved from the WES results and divided into two groups. We observed 1495 and 1033 events in UCKT and UCHD groups. Using the start site as a reference, events occur in two or more patients in either group were selected. We removed records that were found in both groups and only reported those that were specific in UCKT. We identified 208 events with 92 unique CNVs in UCKT group (Additional file 2: Table S1). None of 14 genes of interest in Table 4 were found in this CNV analysis.

### IPA network analysis

Through IPA, we explored the interactions and pathways among GNAQ, SEPTIN6, NTRK3, and IKZF1 (Fig. 4). SEPTIN6 exhibited no connections in the protein network. GNAQ, NTRK3, and IKZF1 were analyzed using the build tool to determine how their relationships are affected by UC. The protein network associated with the three main molecules is involved in the canonical signaling pathways of bladder cancer, PI3K/AKT, and mTOR.
Each molecule was treated as a seed, and the network was constructed by connecting these seeds, thereby enabling us to explore the relationships among them. As indicated in Fig. 4, GNAQ, NTRK3, and IKZF1 may regulate UC through AKT1, MTOR, PI3KR1, HRAS, SRC, TP53, CCDN1, and RUNX3. The relationships among molecules involve activation, causation, correlation, expression, phosphorylation, protein–protein binding, were indicated in the network.

As indicated in Fig. 5, the network analysis revealed that GNAQ, NTRK3, and IKZF1 are related to UC because they are involved in PI3K/AKT and bladder cancer signaling. The 25 interacting proteins in the network (AKT1, CCND1, CDKN1A, CDKN1B, CDKN2A, CXC18, EGFR, ERBB2, FOXO1, HDAC5, HRAS, HSP90AA1, HSP90AB1, MAP2K1, MAPK1, MMP2, MTOR, PIK3CA, PIK3R1, RB1, SIN3A, TP53, TSC2, VEGFA, and RUNX3) warranted further investigation. Thus, the result from IPA analysis explored the target proteins and networks that potentially contribute to UC and bladder cancer signaling pathways, which may inform future studies of the underlying mechanisms of UC.

**Discussion**

Mutational profile analysis is the key to understanding the tumorigenic mechanisms of UC. The mutational profiles of UC that develop after transplantation have not been thoroughly investigated. In this study, WES analysis allowed us to characterize the mutational signatures of UC in detail. The comparison of CNV data in UC and bladder cancer revealed a subset of unique events and was warranted further investigation. Specific nucleotide conversion patterns that develop in certain cancers can elucidate the establishment of these mutations [45, 46]. In a TCGA study, 51% of all the mutations in bladder cancers were TpC > T or G mutations and 3.8% were TpC > A mutations [15]. In addition, TP53, PIK3CA, and FGFR3 are the most common molecular alterations associated with UC [15, 47].
Tumors developed after kidney transplants are believed to be related to various factors, including immunosuppressive agents, viral infections, geographic area, and uremic environment, in patients with CKD/ESKD [48–50]. In the present study, missense mutations were the most common somatic mutations, followed by nonsense mutations. Furthermore, T > A replacements were the most common point mutations. Aristolochic acid (AA), which originates from plants in the genus Aristolochia that were used in herbal medicine, is a powerful nephrotoxin and human carcinogen associated with CKD and upper urinary tract UC (UUC) [51]. Aristolactam–DNA adducts, which are found in urothelial tissues [52], initiate mutation in the TP53 tumor suppressor gene, creating a biomarker specific to AA-induced UUC [52, 53]. Taiwan has an extraordinarily high incidence of UUC due to the widespread use of Aristolochia herbal therapies [54], which suggests that AA might play a central role in the etiology of UUC. The mutation pattern discovered in our patients suggests that a possible history of exposure to Aristolochia remedies led them to develop ESKD and UC.

Four somatic mutations (GNAQ, SEPTIN6, NTRK3, and IKZF1) identified in current study may be related to UCKT. All these mutations were single-base substitutions specifically identified in the UCKT group, and none of the genes had been reported in the genomes of patients with UC. In our study, these mutations were detected in more than one UCKT genome, which suggests they did not occur randomly and potentially played roles in the pathogenesis of UCKT. According to the protein network in IPA, GNAQ, NTRK3, and IKZF1 may regulate
UC through AKT1, MTOR, PI3KR1, HRAS, SRC, TP53, CCND1, and RUNX3. Therefore, PI3K/AKT and bladder cancer signaling and their associated pathways must be further investigated in the future.

G proteins play an essential role in cellular signal transduction. The Gq protein alpha subunit (encoded by GNAQ) couples with a seven-transmembrane receptor to activate phospholipase C-beta, which generates second messengers, and to activate kinase cascades in the cytoplasm of the cells [55]. These signals ultimately control gene transcription and cell survival, motility, and growth. However, signals transmitted by G-protein-coupled receptors/G proteins and downstream targets are involved in the initiation and progression of cancer [56]. GNAQ is located on chromosome 9q21. The role of GNAQ as an oncogene has been observed in uveal melanoma [57], blue nevi [58], and malignancies affecting the meninges [59].

NTRK3 is a member of the NTRK-encoding family, which comprises the genes NTRK1, NTRK2, and NTRK3 that encode TrkA, TrkB, and TrkC, respectively [60]. NTRK3 is located on chromosome 15q25. NTRK receptors signal through the JAK/STAT, PI3K/AKT, and MEK/ERK pathways to promote cell proliferation, differentiation, and survival [61]. Although numerous studies have
investigated the key roles of these receptors in the development and function of the central and peripheral nervous system [62], alterations in NTRK genes have been observed in patients with colon [63], thyroid [64, 65], lung [66], glial [67], and breast [68] cancers. These alterations occur at relatively low frequencies (< 1%) among patients with each of these types of tumors, but NTRK-driven cancers affect numerous patients, making them key therapeutic targets [69–71].

IKZF1 encodes a transcription factor that belongs to the family of zinc finger DNA-binding proteins associated with chromatic remodeling. IKZF1 is located on chromosome 7p12. IKZF1 expression was highly conserved, which suggests its fundamental role in the ontogeny of the lymphopoietic system across species [73]. Ikaros genes are a major determinant of hematopoietic lineage, especially that of lymphocytes [74]. IKZF1 deletions and mutations affect B-cell precursor acute lymphoblastic leukemia and contribute to its poor prognosis [75]. Constitutional and acquired genetic changes in Ikaros genes have been associated with human diseases, including lung, ovarian, liver, and colorectal cancers [74, 76–78].
Cell signaling pathways play key roles in tumorigenesis. Genetic and protein alterations in these pathways can modify cell cycle control, DNA repair, and carcinogen metabolism [79]. The mammalian target of rapamycin (mTOR) pathway is central to the development of multiple cancers, including UC [80]. mTOR is involved in complex signaling cascades that regulate cell growth and angiogenesis under both normal and cancerous conditions. Other key molecules involved in this pathway include upstream activators, such as PI3K; AKT; negative regulators, such as the tuberous sclerosis complex (TSC) 1/2; and downstream effectors, such as p70 S6 kinase and 4EBP1 [81]. The functional role of our candidate genes (GNAQ, NTRK3, and IKZF1) in these pathways identified through IPA require further validation.

This study has several limitations. First, this was a retrospective study with a limited sample size. Second, due to limitations in sample collection, all the patients in our study groups were female; nonetheless, this is consistent with the finding that most UCKT individuals in the Taiwanese population are female [11, 82]. Additional studies with larger sample sizes are required to achieve conclusive results. Third, the contribution of the dbSNPs were not thoroughly discussed. Lastly, functional validation is necessary to identify the characteristics of each of the candidate genes.

Conclusion
In conclusion, we identified 14 novel mutations of CARD11, FNPB1, GNAQ, HOXD13, IKZF1, MAX, MLLT10, NTRK3, SEPTIN6, SEPTIN9, SH3GL1, SLC34A2, TAL1, and TRAF7 in a group of patients with UCKT. Among the affected genes, GNAQ, IKZF1, and NTRK3 were potentially involved in the signaling network of UCKT. These findings could elucidate the development of UCKT and serve as a basis for the discovery of new potential biomarkers and the development of more effective treatments for UCKT.

Abbreviations
ESKD: End-stage kidney disease; PTMs: Post-transplant malignancies; KT: Kidney transplantation; UC: Urothelial carcinoma; TCGA: The Cancer Genome Atlas; UCKT: Urothelial carcinoma post kidney transplantation; WES: Whole-exome sequencing; UCHD: Urothelial carcinoma of hemodialysis patients; CKD: Chronic kidney disease; SNVs or SNPs: Single-nucleotide variations/polymorphisms; InDels: Insertions or deletions; FFPE: Formalin-fixed paraffin-embedded; DNA: Deoxyribonucleic acid; CCG: Cologne Center for Genomics; BWA: Burrows-wheeler Alignment; SBS: Single-base substitution; GATK: Genomic Analysis Toolkit; COSMIC: Catalogue of Somatic Mutations in Cancer; IntOGen: The Integrative Onco Genomic; PCR: Polymerase chain reaction; PBMC: Peripheral blood mononuclear cell; IPA: Ingenuity Pathway Analysis; GNAQ: Guanine nucleotide-binding protein G(q); CNV: Copy number variations; NTRK3: Neurotrophic tyrosine receptor kinase 3; IKZF1: Ikaros family zinc finger protein 1; AA: Anastomotic acid; ULUC: Upper urinary tract urothelial carcinoma; mTOR: Mammalian target of rapamycin.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12967-022-03522-4.

Acknowledgements
The authors acknowledge the Cologne Center for Genomics, University of Cologne, Germany and the Clinical and Industrial Genomic Application Development Service Center of National Core Facility for Biopharmaceuticals, Taiwan (MOST 107-2319-B-010-002) for sequencing. The authors also thank the Center for Research Resources and Development in Kaohsiung Medical University, Kuan-Wen Chen and Tzu-Wei Hung in GGA Corporation, Molecular Science and digital innovation center (Taiwan) for the assistance in performing the QIAGEN IPA analysis. This study was supported by grants from the Kaohsiung Medical University Hospital (KMUH104-4M09, KMUH110-0R18), Taiwan Ministry of Science and Technology (MOST) 109-2635-B-037-009 and MOST 110-2314-B-037-071.

Author contributions
Conceptualization, H.T.K., D.Y.H and L.M.L.; methodology, H.T.K., D.Y.H. and A.M.H.; software, W.Y.C.; formal analysis, W.Y.C, D.Y.H, C.C.Y, P.I.L. and S.M.C; resources, H.L.K.; writing—original draft preparation, L.M.L.; writing—review and editing, L.M.L, W.Y.C, A.M.H and T.W.L; supervision: A.M.H and H.T.K.; All authors have read and agreed to the published version of the manuscript.

Funding
This study was supported by research grants from the Kaohsiung Medical University Hospital (KMUH104-4M09, KMUH110-0R18), Taiwan Ministry of Science and Technology (MOST) 109-2635-B-037-009 and MOST 110-2314-B-037-071.

Availability of data and materials
The data that support the findings presented in this manuscript are available from the corresponding author upon request.

Declarations
Ethics approval and consent to participate
The sample collection and study protocol were conducted under the Institutional Review Board of Kaohsiung Medical University Hospital (KMU-HIRB-G0-20150030). Written informed consent was obtained from all the patients, and all clinical investigations were conducted in accordance with the Declaration of Helsinki.

Consent for publication
The publication of this manuscript has been approved by all authors.

Competing interests
The authors declare no conflicts of interest.

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Received: 11 February 2022  Accepted: 8 July 2022

Published online: 21 July 2022

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