CANCER GENETICS

Identification of Genetic Risk Factors for Familial Urinary Bladder Cancer: An Exome Sequencing Study

Alexander Pemov, PhD1; Talia Wegman-Ostrosky, MD1; Jung Kim, PhD2; Stella Koutros, PhD2; Brenna Douthitt, BA1; Kristine Jones, BSc3; Bin Zhu, PhD1; Dalsu Baris, PhD2; Molly Schwenn, MD2; Alison Johnson, MBA2; Margaret R. Karagas, PhD2; Brian D. Carter, MPH1; Marjorie L. McCullough, ScD, RD1; Maria Teresa Landi, MD, PhD2; Neal D. Freedman, PhD2; Demetrius Albanes, MD1; Debra T. Silverman, ScD2; Nathaniel Rothman, MD, MPH1; Neil E. Caporaso, MD2; Mark H. Greene, MD1; Joseph F. Fraumeni Jr, MD1; and Douglas R. Stewart, MD1

PURPOSE Previous studies have shown an approximately two-fold elevation in the relative risk of urinary bladder cancer (UBC) among people with a family history that could not be entirely explained by shared environmental exposures, thus suggesting a genetic component in its predisposition. Multiple genome-wide association studies and recent gene panel sequencing studies identified several genetic loci that are associated with UBC risk; however, the list of UBC-associated variants and genes is incomplete.

MATERIALS AND METHODS We exome sequenced eight patients from three multiplex UBC pedigrees and a group of 77 unrelated familial UBC cases matched to 241 cancer-free controls. In addition, we examined pathogenic germline variation in 444 candidate genes in 392 The Cancer Genome Atlas UBC cases.

RESULTS In the pedigrees, segregating variants were family-specific although the identified genes clustered in common pathways, most notably DNA repair (MLH1 and MSH2) and cellular metabolism (IDH1 and ME1). In the familial UBC group, the proportion of pathogenic and likely pathogenic variants was significantly higher in cases compared with controls (P = .003). Pathogenic and likely pathogenic variant load was also significantly increased in genes involved in cilia biogenesis (P = .001). In addition, a pathogenic variant in CHEK2 (NM_007194.4: c.1100del; p.T367Mfs*15) was over-represented in cases (variant frequency = 2.6%; 95% CI, 0.71 to 6.52) compared with controls (variant frequency = 0.21%; 95% CI, 0.01 to 1.15), but was not statistically significant.

CONCLUSION These results point to a complex polygenic predisposition to UBC. Despite heterogeneity, the genes cluster in several biologically relevant pathways and processes, for example, DNA repair, cilia biogenesis, and cellular metabolism. Larger studies are required to determine the importance of CHEK2 in UBC etiology.

JCO Precis Oncol 5:1830-1839. © 2021 by American Society of Clinical Oncology

Creative Commons Attribution Non-Commercial No Derivatives 4.0 License

INTRODUCTION
It is estimated that there will be 83,730 newly diagnosed urinary bladder cancers (UBCs: 64,280 males and 19,450 females) and 17,200 deaths (12,260 males and 4,940 females) in the United States in 2021.1 UBC is typically a slowly developing disease but recurs frequently. It is predominantly observed in older patients (average age at diagnosis = 73 years). A number of environmental risk factors have been identified for this malignancy including smoking, some occupational exposures, and contaminants in drinking water.2 Cigarette smoking is the primary risk factor for bladder cancer, which is estimated to account for approximately 50% of UBC cases in both sexes.2,3 Risk among current smokers is four to five times greater than that in nonsmokers.2,3 Besides environmental factors, a genetic component of predisposition to UBC has been demonstrated as well: the first evidence of genetic susceptibility to UBC was observed in a pedigree of four affected first-degree relatives by Fraumeni and Thomas.4 Subsequent epidemiologic studies have identified an increased relative risk for individuals with family history of UBC,5-10 which could not be fully explained by shared environmental exposure, thus implying a genetic component in the predisposition.11-15 However, familial UBC clustering appears to be rare: a national recruitment effort failed to identify a sufficient number of multiple-case UBC kindreds to warrant a familial cancer study.5

Early important clues for a potential UBC genetic etiology came from studies of hereditary cancer susceptibility disorders such as Lynch (eg, OMIM#120435), Costello (OMIM#218040), Apert (OMIM#101200), and familial adenomatous polyposis 3 (OMIM#616415) syndromes. The presence of UBC in these monogenic disorders5 suggests that rare variants in critical signaling pathways (eg, cell cycle progression and mitogenic signal transduction) play an etiologic role in UBC pathogenesis. A subsequent series of pioneering genome-wide association studies

ASSOCIATED CONTENT
Data Supplement
Author affiliations and support information (if applicable) appear at the end of this article.
Accepted on November 3, 2021 and published at ascopubs.org/journal/jp on December 22, 2021; DOI https://doi.org/10.1200/PO.21.00115
has identified 16 common, low-penetration polymorphisms associated with elevated UBC risk.\textsuperscript{2,16-19} A recent genome-wide meta-analysis that investigated the outcomes of non–muscle-invasive UBC identified rs12885353 (near SCFD1), which is significantly associated with UBC recurrence-free survival.\textsuperscript{20} Most of these variants and genes confer modest increase in UBC risk (odds ratio $< 2$) and aggregate in xenobiotic metabolism, DNA repair, and cell cycle progression pathways.\textsuperscript{21}

Two recent studies performed cancer gene panel sequencing in predominantly patients with sporadic UBC. Both investigations identified pathogenic and likely pathogenic (P and LP) variants in germlines of 13.7\%\textsuperscript{22} and 24\%\textsuperscript{23} of patients, with \textit{BRCA1, BRCA2, MSH2, CHEK2, ERCC3, MLH1,} and \textit{ATM} being most frequently mutated. Unlike previous sequencing studies, which analyzed candidate gene panels in primarily sporadic UBC cases\textsuperscript{22,23}, we used an exome-wide approach in patients with familial UBC. We tested the following hypotheses: (1) Familial high-penetration clusters of bladder cancer are partially driven by shared genetic variants that predispose to higher incidence of the cancer cases in related individuals, (2) Rare deleterious variants segregating among multiple cases in each pedigree could be involved in the etiology of familial bladder cancer, and (3) Rare deleterious variants detected by exome sequencing could have effects that are large enough to be detected in a modestly sized sample set. In this exploratory study, we investigated exomes of eight patients from three multiplex UBC pedigrees and 77 unrelated familial UBC cases that were matched to 241 cancer-free controls. In the pedigrees, we ascertained the segregating pattern of deleterious variants, and in the case-control analysis, we examined a rare-variant association with the UBC risk. In addition, we investigated the germline variation landscape in 392 UBC cases from The Cancer Genome Atlas (TCGA) public database (Data Supplement). The sets were analyzed in parallel, and the results were examined for common variants, genes, and pathways.

**MATERIALS AND METHODS**

The full version of Materials and Methods can be found in the Data Supplement.

**Patients and Sample Collection**

All studies were approved by the institutional review board (IRB), the National Cancer Institute (NCI) Special Studies IRB, and participating local IRBs. Clinical information for three pedigrees is summarized in the Data Supplement. Clinical information for 74 familial UBC cases is summarized in the Data Supplement. Cases were matched to controls, and the principal component analysis (PLINK \textit{v1.90b4.4})\textsuperscript{24} was performed on the resulting set to ensure its homogeneity (Data Supplement).

All participants provided written informed consent before enrollment into the NCI DCEG familial cancer protocol "Clinical, Laboratory, and Epidemiologic Characterization of Individuals and Families at High Risk of Cancer" or the parent studies that enrolled the participants. All individual-level data, including clinical data, were deidentified. The authors have modified the pedigree or family tree to avoid potential identification of the family or its members. The authors received and archived written patient consent. This study fully adhered to the principles set out in the Declaration of Helsinki.

**Exome Sequencing and Data Processing**

Genomic DNA was extracted from blood, whole genome amplified (74 familial UBC cases), exome captured with NimbleGen SeqCap EZ Human Exome Library, and sequenced on the Illumina HiSeq 2000 platform. The human reference genome and the known gene transcript annotation were downloaded from the UCSC database, hg19. Sequencing reads were trimmed (Trimmomatic), and only read pairs with both ends $> 36$ bp were used. Reads were aligned to the reference genome (NovoAlign). Duplicate reads were removed (MarkDuplicates), and only read pairs mapped in complementary directions at a fragment length of 200-400 bp were used. These alignments were further refined (RealignerTargetCreator and IndelRealigner). Variant discovery and genotype calling were performed on all individuals globally (UnifiedGenotyper, HaplotypeCaller from GATK, and FreeBayes). The three callers were used to call each sample in parallel, and the caller-specific results were generated independently. The ensemble variant calling pipeline was then implemented to integrate the results from the three callers.

**Data Filtering and Variant Classification**

All noncoding, multiallelic, common variants (> 1\% in ExAC or gnomAD) and variants present in this study’s controls at frequency above 10\% were filtered out. Remaining variants were grouped into three tiers: (1) variants classified in ClinVar as pathogenic or likely pathogenic (tier 1); (2) variants that were unclassified by ClinVar but classified by InterVar as P and LP (tier 2); and (3) all remaining loss-of-function variants and missense variants fulfilling 2 of 3 of the following conditions: CADD\_phred\_score $> 25$, REVEL\_score $> 0.5$, MetaSVM\_score = $D$(eleterious) (tier 3). Variants in tier 1 were considered deleterious; remaining variants (tiers 2 and 3) were considered potentially deleterious.

**Variant Segregation Pattern in Pedigrees**

Tier 1-3 variants found in UBC-affected members of a pedigree were considered as risk variants and were examined further.

**Statistical Tests**

Differences in frequency between cases and controls were determined by using Fisher’s exact test. Rare-variant association tests were performed by using the Cohort Allelic Sums Test, Sequence Kernel Association Test (SKAT), and SKAT optimal test. False discovery rate correction for
multiple testing was computed in variant- and gene-based analyses for case-control association tests (q-value < 0.05). Bonferroni correction was applied to pathway-level analyses (0.05/9 = 0.006, \( P \text{ value}_{\text{corrected}} < .006 \)).

**Ontological Classification of Genes Carrying P and LP Variants**

In the familial UBC case-control analysis, genes with tier 1 P and LP variants were stratified by their biologic processes (BPs) as defined in the Gene Ontology database. Related BP terms were further grouped into the following categories: DNA repair, replication, and recombination, gene expression and signal transduction, cellular metabolism, transmembrane transport, protein modifications and metabolism, and cilia biogenesis. Infrequently observed or biologically irrelevant BP categories were placed in the Others group. Genes with unknown BP were placed in the Unknown group.

**UBC Gene List Compilation**

The list of genes likely involved in the etiology of UBC was compiled by combining genes from published genome-wide association studies, somatic sequencing studies, studies of tumor predisposition syndromes, and all known DNA repair genes (Data Supplement). OncoPrint plots summarize clinical and genomic characteristics for patients carrying tier 1-2 variants in the resulting 444 candidate genes.

**TCGA UBC Data Set**

Germline sequencing data for UBC-diagnosed participants (\( N = 392 \)) were downloaded from the Genomic Data Commons. Common variants (> 1%) were filtered out. Tier 1 and 2 variants were used for further analysis.

**RESULTS**

**Variant Segregation Pattern in Three Multiplex UBC Pedigrees**

The UBC pedigrees analyzed in this study are shown in Figure 1. Clinical information for these families is summarized in the Data Supplement.

We exome sequenced germline DNA from three, two, and three UBC-affected members of families A, B, and C, respectively. After ascertainment of variant segregation pattern and assigning the variants to tiers 1, 2, and 3, we identified 4, 15, and 12 tier 1-3 variants in the pedigrees A, B, and C, respectively (Table 1). We detected a single tier 2 variant (P and LP InterVar) in \( \text{CFTR} \) in family A, two tier 2 variants (\( \text{IDH1} \) and \( \text{ELAC2} \)) in family B, and one tier 1 (\( \text{ABCA4} \)) and one tier 2 (\( \text{CHRNE} \)) variants in family C. rs119484086 in \( \text{ELAC2} \) has been reported as a prostate cancer susceptibility allele\(^{85}\); however, there were no cases of prostate cancer reported for the members of family B who harbored the variant. \( \text{ABCA4} \), \( \text{CHRNE} \), and \( \text{CFTR} \) are expressed at a low level in the bladder, and their known biologic functions (retina-specific membrane transporter, acetylcholine receptor at neuromuscular junctions, and water secretion and absorption in epithelial tissues, respectively) make them candidates unlikely for UBC predisposition. In families B and C, we identified tier 3 variants in mismatch repair genes \( \text{MLH1} \) and \( \text{MSH2} \), respectively. In addition to a tier 2 variant segregating in family B in \( \text{IDH1} \) (one of the key enzymes of carbon metabolism in the cell), we identified a tier 3 variant in \( \text{ME1} \) (malic enzyme 1, which connects the glycolytic pathway with the Krebs cycle) that segregated in family A.

**Exome-Wide Analysis of 77 Familial UBC Cases Versus 241 Cancer-Free Controls**

**Variant-level analysis by Fisher’s exact test.** We observed only one single variant in \( \text{ATP2A1} \) that reached statistical significance after multiple testing correction (q-value < 0.05; Table 2). \( \text{ATP2A1} \) is unexpressed in the urinary bladder and was not investigated further. Notably, the frequency of frameshifting deletion in \( \text{CHEK2} \) (NM_007194:c.1100del;p.T367Mfs*15, rs555607708) was elevated among cases (2.6%; 95% CI, 0.71 to 6.52) compared with controls (0.21%; 95% CI, 0.0 to 1.15) by approximately 10-fold; however, it was not significant after multiple testing correction. In several largest public databases, the frequency of c.1100delC among Europeans (excluding Finnish subpopulation) ranged from 0.17% to 0.26%; its frequency varied between different ancestral groups and was highest among Finns (0.87%; 95% CI, 0.76 to 0.99; Table 3).

**Gene-level analysis by Cohort Allelic Sums Test, SKAT, and SKAT optimal test rare-variant association (burden) tests.** The gene-level analysis identified \( \text{CC2D2A} \) and \( \text{GALC} \) at the nominal 0.05 significance level by at least one of the tests, but neither of these genes were significant after multiple testing correction (Data Supplement).

**Comparison of P and LP variant loads in 77 cases versus 241 controls.** In addition to variant- and gene-level analyses, we examined the load of tier 1 (ClinVar P and LP) variants in 77 cases and 241 controls. Visual inspection of the distribution of the number of P and LP variants per person in cases and controls revealed a shift toward a higher number of P and LP variants in cases (Fig 2A). We also observed a higher proportion of individuals with at least one P and LP variant among cases as compared with controls (76.6% vs 66.4%, \( P = .003 \), Table 4). This difference was statistically significant after Bonferroni correction. The total number of unique and overlapping P and LP variants and genes in cases and controls is shown in Figure 2B.

**Ontological analysis of tier 1 P and LP variants in 77 cases and 241 controls.** First, we stratified genes harboring P and LP variants into groups with related BP as defined in the Gene Ontology database (Data Supplement). We then determined the proportion of individuals who carried at least one P and LP variant in any of the genes included in an ontological category as referenced above, in both cases and controls. The cases versus controls comparison
demonstrated that for most of the ontological categories, the proportion of individuals with at least one P and LP variant was higher among cases; however, after multiple testing correction (Bonferroni), the differences reached statistical significance only in the cilia biogenesis category ($P = .001$; Table 4 and Figs 2C and 2D).

**Analysis of Pathogenic Variant Loads in the Germline of 392 UBC Cases From TCGA Identified an Elevated Frequency of CHEK2 c.1100delC**

We also examined ClinVar and InterVar P and LP variants in 444 UBC candidate genes (Data Supplement) found in the germlines of 392 TCGA UBC patients (Data Supplement). In total, we observed 123 tier 1 and 2 variants in 59 genes among TCGA UBC patients. Variants in CHEK2 were observed in 9 of 392 (2.3%) patients, thus making this locus the most frequently altered among 444 candidate genes in the UBC TCGA set (1.15%; 95% CI, 0.53 to 2.17). Notably, 4 of 9 (44.4%) of these CHEK2 pathogenic variants were the deletion c.1100delC identified in the familial UBC group; its frequency among 392 UBC cases was 0.51% (95% CI, 0.14 to 1.30; Table 3).

**DISCUSSION**

In this exploratory study, we investigated genetic risk factors in familial UBC using exome sequencing data from three multiplex pedigrees and 77 familial UBC cases matched with 241 cancer-free controls from existing epidemiologic studies and examined pathogenic germline variant loads in 444 UBC candidate genes in 392 UBC cases from the TCGA set. In the pedigrees, we identified potentially deleterious variants in mismatch repair DNA repair genes MLH1 and MSH2 that segregated in families B and C, and in the carbon metabolism genes, IDH1 and ME1, in families B and A. In the analysis of the familial UBC cases versus controls, we identified a possible association between the CHEK2 c.1100delC pathogenic variant and UBC, and in the TCGA UBC set, we observed this CHEK2 pathogenic variant at somewhat elevated frequency as well (0.51%; 95% CI, 0.14 to 1.30). Finally, we found that cilia biogenesis genes were significantly enriched with P and LP variants and that the total P and LP variant load was significantly higher in 77 cases with a positive UBC family history compared with controls from the epidemiologic studies. The main limitation of this study was a modest number of samples. This obstacle, which is common in projects involving rare diseases such as familial UBC, precluded us from reaching a sufficient power despite the extensive effort. Future replication studies would benefit from broad collaborations.

The variant segregation pattern in the three pedigrees demonstrated that the variants and the variant-carrying genes were unique to each family. Yet, we identified common ontological categories and biologic pathways...
| Family ID | Gene Name | Chr (hg19) | Variant ID | Reference Allele | Variant Allele | Type of Variant Allele | gnomAD Population Allele Frequency | ClinVar Call | InterVar Call | MetaSVM Prediction | CADD Score | REVEL Score | Expression in Urinary Bladder |
|-----------|------------|------------|------------|------------------|----------------|------------------------|-------------------------------------|--------------|---------------|---------------------|-----------|-------------|-----------------------------|
| A         | CFTR       | 7          | rs397508137 | G                | A              | Missense               | 0.0001                              | VUS          | LP            | D                   | 19.8      | 0.692       | Low, 1                      |
| A         | STEAP3     | 2          | rs199836424 | G                | A              | Missense               | 0.002                               | —            | LB            | D                   | 29.7      | 0.860       | Medium, 6                   |
| A         | DHA5H      | 5          | rs78853309  | C                | G              | Missense               | 0.0002                              | VUS          | VUS          | T                   | 26.5      | 0.688       | Low, 1                      |
| A         | ME1        | 6          | rs375470975 | A                | C              | Missense               | 0.00005                             | VUS          | T             | —                   | 27.7      | 0.646       | Medium, 5                   |
| B         | IDH1       | 2          | rs762820641 | C                | T              | Missense               | 0.00001                             | —            | LP            | D                   | 25.9      | 0.779       | High, 97                    |
| B         | ELAC2      | 17         | rs118484086 | C                | T              | Missense               | 0.00005                             | —            | LP            | D                   | 35.0      | 0.491       | Medium, 10                  |
| B         | PADI3      | 1          | rs199619967 | C                | T              | Stopgain              | 0.0002                              | —            | —             | —                   | 28.3      | —           | Medium, 7                   |
| B         | ASIC4      | 2          | rs20379222  | C                | T              | Stopgain              | —                                   | —            | T             | —                   | 36.0      | —           | Unexpressed                 |
| B         | SLC13A1    | 7          | rs28364172  | G                | A              | Stopgain              | 0.002                               | VUS          | —             | —                   | 34.0      | —           | Unexpressed                 |
| B         | RGL4       | 22         | rs748038406 | A                | AC             | Frameshift            | —                                   | VUS          | —             | —                   | 24.1      | —           | Low, 3                      |
| B         | MLH1       | 3          | rs35052531  | AA               | GC             | Delins                | 0.003                               | LB D B       | 27.8          | 0.659-0.963b      | Medium, 5 |
| B         | NOC21      | 1          | rs143094540 | C                | T              | Missense               | 0.0001                              | VUS          | T             | —                   | 28.4      | 0.599       | Low, 12                     |
| B         | DOCK3      | 3          | 51411957    | C                | T              | Missense               | —                                   | VUS          | D             | —                   | 34.0      | 0.562       | Unexpressed                 |
| B         | CACNA1D    | 3          | rs763788750 | A                | G              | Missense               | 0.00002                             | VUS          | D             | —                   | 24.7      | 0.612       | Low, 0.4                    |
| B         | AEBP1      | 7          | rs370857030 | A                | G              | Missense               | 0.00002                             | —            | VUS          | —                   | 24.2      | 0.769       | Medium, 50                  |
| B         | CIS        | 12         | 7177845     | G                | T              | Missense               | 0.000004                            | VUS          | D             | —                   | 26.3      | 0.771       | High, 245                   |
| B         | SLC18B7    | 12         | rs60786449  | C                | T              | Missense               | 0.00001                             | VUS          | D             | —                   | 25.0      | 0.543       | Unexpressed                 |
| B         | NFSE1      | 20         | rs11246981  | T                | C              | Missense               | 0.004                               | —            | VUS          | D                   | 20.3      | 0.656       | Low, 4                      |
| B         | SCN9A      | 2          | rs141268327 | T                | C              | Missense               | 0.004                               | LB LB D       | 23.9          | 0.899       | Low, 0.2                   |
| B         | ABCA4      | 1          | rs1750120   | G                | A              | Missense               | 0.00032                             | P LP D       | 35.0          | 0.887       | Low, 0.2                   |
| B         | CHRNE      | 17         | rs14002330  | C                | T              | Missense               | 0.0007                              | VUS          | LP            | D                   | 27.0      | 0.681       | Low, 1                      |
| B         | ITGAV      | 2          | 187529302   | C                | A              | Stopgain              | —                                   | VUS          | —             | —                   | 35.0      | —           | High, 23                    |
| B         | LRRF1P2    | 3          | rs14960210  | G                | A              | Stopgain              | 0.0009                              | VUS          | —             | —                   | 39.0      | —           | Medium, 6                   |
| B         | IGSF10     | 3          | rs14296318  | G                | A              | Stopgain              | 0.0007                              | VUS          | —             | —                   | 36.0      | —           | Low, 2                      |
| C         | VWA2       | 10         | 11632630    | TG               | T              | Frameshift            | —                                   | VUS          | —             | —                   | 23.5      | —           | Low, 0.4                    |
| C         | MSH2       | 2          | rs587779113 | A                | C              | Missense               | 0.00004                             | VUS          | D             | —                   | 23.2      | 0.720       | Low, 2                      |
| C         | RABL2A     | 2          | rs145167719 | G                | A              | Missense               | 0.004                               | —            | VUS          | T                   | 25.6      | 0.686       | Low, 2                      |
| C         | NEAL2      | 3          | rs203173710 | G                | A              | Missense               | 0.002                               | VUS B T       | 32.0          | 0.610       | Medium, 5                   |
| C         | SLC6A19    | 5          | rs7629880   | C                | T              | Missense               | 0.0003                              | —            | VUS          | D                   | 28.9      | 0.877       | Unexpressed                 |
| C         | PSD2       | 5          | rs138380367 | G                | A              | Missense               | 0.0007                              | VUS          | D             | —                   | 28.5      | 0.819       | Unexpressed                 |
| C         | MYO18A     | 17         | rs76590796  | C                | T              | Missense               | 0.003                               | VUS          | D             | —                   | 35.0      | 0.488       | Medium, 7                   |

Abbreviations: —, data not available; B, benign; D, disease-causing; delins, deletion-insertion; LB, likely benign; LP, likely pathogenic; P, pathogenic; T, tolerated; VUS, variant of unknown significance.

*Expression in urinary bladder: gene expression values were obtained from NCBI Gene db25 and are shown in RPKM (Reads Per Kilobase of transcript, per Million mapped reads) units.

*Range of REVEL scores for possible substitutions of MLH1 K618 for N, R, Q, M, E, and T amino acids.
affected by these variants in the pedigrees. For instance, we observed potentially deleterious variants in \textit{MSH2} and \textit{MLH1} segregating in families C and B, respectively. A rare missense \textit{MSH2} variant (c.182A>C; p.Q61P) found in family C was also identified in a patient who fulfilled the Bethesda guidelines for Lynch syndrome and who developed an ovarian cancer and colorectal carcinoma at age 44 and 50 years, suggesting a causative role of this variant.\textsuperscript{27} A deletion-insertion \textit{MLH1} variant (rs35502531)\textsuperscript{28} segregating in family B (c.1852_1853delinsGC; p.K618A), although classified as benign by ClinVar, has been shown to weaken the interaction between MLH1 and PMS2 in functional studies.\textsuperscript{29}

We also observed rare deleterious and potentially deleterious variants in the carbon metabolism genes, \textit{IDH1} and \textit{ME1}, in Families B and A, respectively. The enzymatic activity of \textit{IDH1} and \textit{ME1} results in increased cellular concentration of nicotinamide adenine dinucleotide phosphate, reduced (NADPH), which could be used to neutralize the excess of reactive oxygen species produced by stress stimuli including xenobiotics.\textsuperscript{30,31} It should be mentioned that in pedigrees A and B, 11 of 13 patients with UBC were current or former smokers. One possible nexus between mutants \textit{ME1} and \textit{IDH1} in the etiology of smoking-related UBC could be a consequence of decreased efficiency of these two enzymes in detoxicating xenobiotics produced by tobacco use.

The frameshift deletion (c.1100del; p.T367Mfs*15) in \textit{CHEK2} was one of the most frequently observed pathogenic variants in this study. \textit{CHEK2} is a serine-threonine kinase that regulates DNA repair through phosphorylation of BRCA2 and arrests progression through the cell cycle via DNA double-strand breaks activation pathway.\textsuperscript{32} The c.1100delC variant has been shown to eliminate kinase activity of \textit{CHEK2} and increase risk of breast cancer 2-fold in women and 10-fold in men.\textsuperscript{33} In ClinVar, this variant is

### TABLE 2. Fisher’s Exact Test of Association in the Set of 77 Urinary Bladder Cancer Cases Versus 241 Cancer-Free Controls

| Gene Name | Chr | Position (hg19) | Variant ID | Reference Allele | Variant Allele | Type of Variant Allele | gnomAD Population Allele Frequency | ClinVar Call | Inheritance Mode | No. of Cases With Variant | No. of Cases Without Variant | No. of Controls With Variant | No. of Controls Without Variant | Fisher’s Exact Test P | FDR q value |
|-----------|-----|-----------------|------------|------------------|----------------|------------------------|-----------------------------------|-------------|------------------|--------------------------|----------------------------|-----------------------------|-------------------------------|-----------------------------|--------------------------|
| ATP2A1    | 16  | 28913639        | rs751365374| G               | GC             | Frameshift            | 0.0003                            | P           | AR               | 7                        | 70                         | 1                           | 240                          | .0003                      | 0.02                     |
| CHEK2     | 22  | 29091856        | rs555607708| AG              | A              | Frameshift            | 0.0025                            | P           | AD               | 4                        | 73                         | 1                           | 240                          | .013                       | 0.7                      |
| SERPINC1  | 1   | 173883881       | rs121969551| G               | A              | Missense              | 0.0009                            | P           | AD or AR         | 3                        | 74                         | 2                           | 241                          | .014                       | 0.7                      |
| ZMPSTE24  | 1   | 40756542        | rs555339565| G               | GT             | Frameshift            | 0.0006                            | P           | AR               | 2                        | 75                         | 0                           | 241                          | .06                       | 0.9                      |
| ABC4      | 1   | 94508323        | rs61750120 | G               | A              | Missense              | 0.0002                            | P           | AR               | 2                        | 75                         | 0                           | 241                          | .06                       | 0.9                      |
| CRB1      | 1   | 197297973       | rs748136623| G               | GAGTAAT        | In_frame             | 0.001                            | P           | AD or AR         | 2                        | 75                         | 0                           | 241                          | .06                       | 0.9                      |
| CUBN      | 10  | 16960686        | rs757649673| A               | ATACCTC        | Frameshift            | 0.0003                            | P           | AR               | 2                        | 75                         | 0                           | 241                          | .06                       | 0.9                      |
| GNRHR     | 4   | 68619793        | rs10489383| T               | C              | Synonymous            | 0.0041                            | P           | AR               | 3                        | 74                         | 2                           | 239                          | .09                       | 0.9                      |

NOTE. Statistically significant variant (q value < 0.05) is shown in bold font. Abbreviations: AD, autosomal dominant; AR, autosomal recessive; FDR, false discovery rate; P, pathogenic.

### TABLE 3. Frequency of CHEK2 c.1100delC Allele in Sample Sets Used in This Study and in Unaffected Populations

| Study Name | Population | No. of All Alleles | Reference Allele | Variant Allele | No. of Variant Alleles | Variant Allele Frequency (%) | 95% CI (%) |
|------------|------------|---------------------|------------------|----------------|------------------------|----------------------------|------------|
| This study, 77 UBC cases | European | 154 | C | delC | 4 | 2.60 | 0.71 to 6.52 |
| This study, 241 controls | European | 482 | C | delC | 1 | 0.21 | 0.01 to 1.15 |
| TCGA UBC cases | All | 784 | C | delC | 4 | 0.51 | 0.14 to 1.30 |
| Nassar et al, 2019,a UBC cases | All | 1724 | C | delC | 3 | 0.17 | 0.04 to 0.51 |
| ExAC | All | 118,290 | C | delC | 215 | 0.18 | 0.16 to 0.21 |
| ExAC | European_Fin | 6,608 | C | delC | 54 | 0.82 | 0.61 to 1.07 |
| ExAC | European_Non-Fin | 64,922 | C | delC | 152 | 0.23 | 0.20 to 0.27 |
| gnomAD | All | 280,390 | C | delC | 591 | 0.21 | 0.19 to 0.23 |
| gnomAD | European_Fin | 25,124 | C | delC | 219 | 0.87 | 0.76 to 0.99 |
| gnomAD | European_Non-Fin | 127,908 | C | delC | 327 | 0.26 | 0.23 to 0.28 |
| NHLBI ESP | All | 12,504 | C | delC | 15 | 0.12 | 0.07 to 0.20 |
| NHLBI ESP | European American | 8,248 | C | delC | 14 | 0.17 | 0.09 to 0.28 |

Abbreviations: ESP, Exome Sequencing Project; Fin, Finnish population; NHLBI, National Heart, Lung, and Blood Institute; Non-Fin, Non-Finnish population; TCGA, The Cancer Genome Atlas; UBC, urinary bladder cancer.

aNassar et al: Prevalence of pathogenic germline cancer risk variants in high-risk urothelial carcinoma. \textit{Genet Med}, 2019.
classified as pathogenic in 37 reports, as a variant of unknown significance in two, and as a risk factor for breast, colorectal, and prostate cancers in another three submissions.\(^3^4\) Despite its apparent pathogenicity, this variant is relatively common in the general population: its global frequency in gnomAD is 0.21% (95% CI, 0.19 to 0.23) and it fluctuates widely in subpopulations and is the highest in Finnish Europeans (0.87%; 95% CI, 0.76 to 0.99).\(^3^5\) Notably, in our study, we observed this variant at substantially increased frequency (2.6%; 95% CI, 0.71 to 6.52) among 77 UBC cases of European descent. We also found this variant at somewhat elevated frequency (0.51%; 95% CI, 0.14 to 1.30) among 392 TCGA UBC cases. Contrary to our findings, a recent study by Nassar et al reported the frequency of this variant to be equal to 0.17% (95% CI, 0.04 to 0.51) in their set of UBC samples\(^3^5\); however, their cases included a substantial proportion of non-European samples, which could be a contributing factor to the differences observed in the outcomes. Despite its established role in breast and testicular cancers,\(^3^3\),\(^3^6^-^3^8\) no significant association between UBC and c.1100delC has been reported to date. In the Copenhagen general population study, which investigated association of CHEK2 c.1100delC with the risk of breast and other cancers, including UBC, the authors reported a modestly increased hazard ratio of 2.26 (95% CI, 0.94 to 5.43) for UBC, which notwithstanding was nonsignificant (\(P = .07\)).\(^3^9\) Another case-control study from Poland compared combined frequency of four pathogenic founder CHEK2 variants, including c.1100delC, and observed a modestly increased but statistically significant odds ratio of 1.9 (95% CI, 1.3 to 2.7; \(P = .0003\)).\(^4^0\) Substantially larger studies are needed to
TABLE 4. Comparison of P and LP Variant Loads in Different Ontological Categories in 77 Urinary Bladder Cancer Cases Versus 241 Cancer-Free Controls

| Ontological Category                     | Proportion of Cases With at Least One P and LP Variant (%) | Proportion of Controls With at Least One P and LP Variant (%) | Fisher’s Exact Test P |
|-----------------------------------------|------------------------------------------------------------|---------------------------------------------------------------|----------------------|
| DNA repair, replication, and recombination | 16.9                                                      | 10.8                                                         | .17                  |
| Gene expression and signal transduction | 14.3                                                      | 10.8                                                         | .42                  |
| Cellular metabolisma                      | 41.6                                                      | 28.2                                                         | .013                 |
| Transmembrane transport                   | 24.7                                                      | 19.9                                                         | .42                  |
| Protein modifications and metabolism      | 16.9                                                      | 12.9                                                         | .45                  |
| Cilia biogenesis                          | 7.8                                                       | 0.4                                                          | .001                 |
| Others                                   | 13.0                                                      | 14.9                                                         | .85                  |
| Unknown                                  | 7.8                                                       | 8.3                                                          | 1                    |
| All categories, combinedb                 | 76.6                                                      | 66.4                                                         | .003                 |

NOTE. Statistically significant ontological categories (Bonferroni correction 0.05/9 = 0.006, P value < .006) are shown in bold font.

Abbreviation: P and LP, pathogenic and likely pathogenic.

*When appropriate, the chi-squared test was performed instead of Fisher’s exact test.

estimate penetrance of CHEK2 deleterious variants in various subpopulations and to determine this kinase’s role in UBC pathogenesis.

Among other DNA damage repair genes, we observed P and LP variants in BRCA2, ATM, CHEK2, BRIP1, and MUTYH in 16.9% of cases in our familial UBC group. Similar to our findings, two recent papers reported P and LP variants in highly penetrant DNA repair genes in 11.3% and 16.7% of patients with sporadic high-risk UBC. However, in our familial UBC group, we detected P and LP variants only in moderately penetrant genes (except for BRCA2) such as CHEK2, ATM, BRIP1, and MUTYH, whereas highly penetrant genes were variant-free. This difference may be due to the advanced stage and grade of UBC cases analyzed in the abovementioned reports, whereas most of our cases were predominantly (62%) non–muscle-invasive, stage < T2 tumors.

In the ontological analysis of variants and genes overrepresented in cases in our familial UBC group, cilia biogenesis was the only statistically significant category: 7.8% of cases had at least one deleterious variant (CC2D2A, DNAAF4, DNAH5, IQCB1, and RSPH1) versus 0.4% controls (NPHP3; P = .001). There is rapidly accumulating evidence of cilia’s involvement in cancer development and progression.41–43 Interestingly, rs8173 in AURKA (involved in regulation of cilia disassembly in mitosis) conferred significantly greater susceptibility to bladder cancer.44

In conclusion, analyses of three distinct data sets revealed multiple biologically plausible genes that may be associated with UBC etiology, pointing to a complex polygenic character of genetic predisposition to this malignancy. Nonetheless, despite the substantial heterogeneity among these genes, they clustered in a limited number of BP, most notably DNA repair, cilia biogenesis, and cellular metabolism.
AUTHORS’ DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

The following represents disclosure information provided by authors of this manuscript. All relationships are considered compensated unless otherwise noted. Relationships are self-held unless noted. I = Immediate Family Member, Inst = My Institution. Relationships may not relate to the subject matter of this manuscript. For more information about ASCO’s conflict of interest policy, please refer to www.asco.org/rcw or ascopubs.org/go/authors/author-center.

Open Payments is a public database containing information reported by companies about payments made to US-licensed physicians (Open Payments).

REFERENCES

1. Siegel RL, Miller KD, Fuchs HE, et al: Cancer statistics, 2021. CA Cancer J Clin 71:7-33, 2021
2. Silverman DT, Kourous S, Figuerda JD, et al, in Thun MJ, Linet MS, Cerhan JR, et al (ed): Schottenfeld and Fraumeni Cancer Epidemiology and Prevention (ed 4). New York, NY, Oxford University Press, 2018, pp 977-996
3. Freedman ND, Silverman DT, Hollebeck AR, et al: Association between smoking and risk of bladder cancer among men and women. JAMA 306:737-745, 2011
4. Fraumeni JF Jr, Thomas LB: Malignant bladder tumors in a man and his three sons. JAMA 201:507-509, 1967
5. Mueller CM, Caporaso N, Greene MH: Familial and genetic risk of transitional cell carcinoma of the urinary tract. Urol Oncol 26:451-464, 2008
6. Kantor AF, Hartege P, Hoover RN, et al: Familial and environmental interactions in bladder cancer risk. Int J Cancer 35:703-706, 1985
7. Aben KK, Witjes JA, Schoenborg MP, et al: Familial aggregation of urothelial cell carcinoma. Int J Cancer 98:274-278, 2002
8. Murta-Nascimento C, Silverman DT, Kogevinas M, et al: Risk of bladder cancer associated with family history of cancer: Do low-penetrance polymorphisms account for the increase in risk? Cancer Epidemiol Biomarkers Prev 16:1595-1600, 2007
9. Pina K, Hemminki K: Familial bladder cancer in the National Swedish Family Cancer Database. J Urol 166:2129-2133, 2001
10. Randi G, Pelucchi C, Negri E, et al: Family history of urogenital cancers in patients with bladder, renal cell and prostate cancers. Int J Cancer 121:2748-2752, 2007
11. Czenk Lichtenstein P, Hemminki K: Environmental and heritable causes of cancer among 9.6 million individuals in the Swedish Family-Cancer Database. Int J Cancer 99:260-266, 2002
12. Lichtenstein P, Holm NV, Verkasalo PK, et al: Environmental and heritable factors in the causation of cancer—Analyses of cohorts of twins from Sweden, Denmark, and Finland. N Engl J Med 343:78-85, 2000
13. Dong C, Hemminki K: Modification of cancer risks in offspring by sibling and parental cancers from 2,112,616 nuclear families. Int J Cancer 92:214-150, 2001
14. Martin C, Leiser CL, O’Neill B, et al: Familial cancer clustering in urothelial cancer: A population-based case-control study. J Natl Cancer Inst 110:527-533, 2018
15. Kramer AA, Graham S, Burnett WS, et al: Familial aggregation of bladder cancer stratified by smoking status. Epidemiology 2:145-148, 1991
16. Kienemen LA, Sulem P, Besenbacher S, et al: A sequence variant at 4q16.3 confers susceptibility to urinary bladder cancer. Nat Genet 42:415-419, 2010
17. Kienemen LA, Thorlacius S, Sulem P, et al: Sequence variant on 8q24 confers susceptibility to urinary bladder cancer. Nat Genet 40:1307-1312, 2008
18. Rothman N, Garcia-Donuas M, Chatterjee N, et al: A multi-stage genome-wide association study of bladder cancer identifies multiple susceptibility loci. Nat Genet 49:978-984, 2010
19. Wu X, Ye Y, Kienemen LA et al: Genetic variation in the prostate stem cell antigen gene PSCA confers susceptibility to urinary bladder cancer. Nat Genet 41:991-995, 2009
20. Galesloot TE, Grottenhuis AJ, Kolev D, et al: Genome-wide meta-analysis identifies novel genes associated with recurrence and progression in non-muscle-invasive bladder cancer. Eur Urol Oncol. 10.1016/j.euo.2021.07.001 [epub ahead of print on August 2, 2021]
21. de Maturana EL, Rava M, Anumudu C, et al: Bladder cancer genetic susceptibility: A systematic review. Bladder Cancer 4:215-226, 2018
22. Carlo MI, Ravichandran V, Srinarvasan P, et al: Cancer susceptibility mutations in patients with urothelial malignancies. J Clin Oncol 38:406-414, 2020
23. Nassar AH, Abou Alaiwi S, AlDubayan SH, et al: Prevalence of pathogenic germline cancer risk variants in high-risk urothelial carcinoma. Genet Med 22:709-718, 2020
24. PLINK 1.9 home: https://www.cog-genomics.org/plink/1.9/
25. NCBI Gene: https://www.ncbi.nlm.nih.gov/gene
26. NCBI ClinVar: https://www.ncbi.nlm.nih.gov/clinvar/variation/5058/evidence/
27. Loizidou MA, Neophytou I, Papamichael D, et al: The mutational spectrum of Lynch syndrome in Cyprus. PLoS One 9:e105501, 2014
28. NCBI dbSNP: https://www.ncbi.nlm.nih.gov/snp/rs35502531
29. Andersen SD, Liberti SE, Lutzen A, et al: Functional characterization of MLH1 missense variants identified in Lynch syndrome patients. Hum Mutat 33:1647-1655, 2012
30. Gagné F: Chapter 6—Oxidative stress, in Gagné F (ed): Biochemical Ecotoxicology. Oxford, United Kingdom, Academic Press, 2014, pp 103-115
31. Vander Heiden MG, DeBerardinis RJ: Understanding the intersections between metabolism and cancer biology. Cell 168:657-669, 2017
32. UniProt Knowledgebase: https://www.uniprot.org/uniprot/O96017
33. Meijers-Heijboer H, van den Ouweland A, Klijn J, et al: Low-penetrance susceptibility to breast cancer due to CHEK2(*1100delC in noncarriers of BRCA1 or BRCA2 mutations. Nat Genet 31:55-59, 2002
34. NCBI ClinVar: https://www.ncbi.nlm.nih.gov/clinvar/variation/128042/
35. Genome Aggregation Database: https://gnomad.broadinstitute.org/variant/22-29091856-AG-A?dataset=gnomad_r2_1
36. Liang M, Zhang Y, Sun C, et al: Association between CHEK2*1100delC and breast cancer: A systematic review and meta-analysis. Mol Diagn Ther 22:397-407, 2018
37. Vahteristo P, Bartkova J, Eerola H, et al: A CHEK2 genetic variant contributing to a substantial fraction of familial breast cancer. Am J Hum Genet 71:432-438, 2002
38. AlDubayan SH, Pyle LC,Gamulin M, et al: Association of inherited pathogenic variants in checkpoint kinase 2 (CHEK2) with susceptibility to testicular germ cell tumors. JAMA Oncol 5:514-522, 2019
39. Näsland-Koch C, Nordsgaard BG, Bojesen SE: Increased risk for other cancers in addition to breast cancer for CHEK2*1100delC heterozygotes estimated from the copenhagen general population study. J Clin Oncol 34:1208-1216, 2016
40. Żłowocka E, Cybulski C, Górska B, et al: Germline mutations in the CHEK2 kinase gene are associated with an increased risk of bladder cancer. Int J Cancer 122:583-586, 2008
41. Fabbri L, Bost F, Mazure NM: Primary cilia in cancer hallmarks. Int J Mol Sci 20:1336, 2019
42. Higgins M, Obaidi I, Mc Morrow T: Primary cilia and their role in cancer. Oncol Lett 17:3041-3047, 2019
43. Liu H, Kiseleva AA, Golemis EA: Ciliary signalling in cancer. Nat Rev Cancer 18:511-524, 2018
44. Andrew AS, Gui J, Sanderson AC, et al: Bladder cancer SNP panel predicts susceptibility and survival. Hum Genet 125:527-539, 2009