Comprehensive Molecular Characterization of Urothelial Bladder Carcinoma

The Cancer Genome Atlas Research Network

Abstract

Urothelial carcinoma of the bladder is a common malignancy that causes approximately 150,000 deaths per year worldwide. To date, no molecularly targeted agents have been approved for the disease. As part of The Cancer Genome Atlas project, we report here an integrated analysis of 131 urothelial carcinomas to provide a comprehensive landscape of molecular alterations. There were...
statistically significant recurrent mutations in 32 genes, including multiple genes involved in cell cycle regulation, chromatin regulation, and kinase signaling pathways, as well as 9 genes not previously reported as significantly mutated in any cancer. RNA sequencing revealed four expression subtypes, two of which (papillary-like and basal/squamous-like) were also evident in miRNA sequencing and protein data. Whole-genome and RNA sequencing identified recurrent in-frame activating FGFR3-TACC3 fusions and expression or integration of several viruses (including HPV16) that are associated with gene inactivation. Our analyses identified potential therapeutic targets in 69% of the tumours, including 42% with targets in the PI3K/AKT/mTOR pathway and 45% with targets (including ERBB2) in the RTK/MAPK pathway. Chromatin regulatory genes were more frequently mutated in urothelial carcinoma than in any common cancer studied to date, suggesting the future possibility of targeted therapy for chromatin abnormalities.

Introduction

Urothelial carcinoma of the bladder is a major cause of morbidity and mortality worldwide, causing an estimated 150,000 deaths per year\(^1\). Previous studies have identified multiple regions of somatic copy number alteration, including amplification of \(PPARG, E2F3, EGFR, CCND1\) and \(MDM2\), as well as loss of \(CDKN2A\) and \(RB1\)\(^2,3\). Sequencing of candidate pathways has identified recurrent mutations in \(TP53, FGFR3, PIK3CA, TSC1, RB1\) and \(HRAS\)\(^2,3\). More recently, a candidate gene study identified mutations at >10% frequency in several chromatin remodeling genes: \(KDM6A, CREBBP, EP300\), and \(ARID1A\)\(^4\). Focused molecular analyses\(^5,6\) have delineated tumour subtypes and identified kinase-activating FGFR3 gene fusions\(^7,8\).

We report here a comprehensive, integrated study of 131 high-grade muscle-invasive urothelial bladder carcinomas as part of The Cancer Genome Atlas (TCGA) project. Included are data on DNA copy number, somatic mutation, mRNA and microRNA expression, protein and phosphorylated protein expression, DNA methylation, transcript splice variation, gene fusion, viral integration, pathway perturbation, clinical correlates and histopathology to characterize the molecular landscape of urothelial carcinoma. This study identifies a number of mutations and regions of copy number variation that involve genes not previously reported as altered in a significant fraction of bladder cancers. It also identifies potential therapeutic targets in most of the samples analyzed.

Results

Demographic, clinical and pathological data

Samples, (from 19 tissue source sites), consisted of 131 chemotherapy-naïve, muscle-invasive, high-grade urothelial tumours (T2-T4a, Nx, Mx), as well as peripheral blood (n=118) and/or tumour-adjacent, histologically normal-appearing bladder tissue (n=23). Cases were retained only if they met the following criteria: tumour nuclei constituted ≥60% of all nuclei, tumour necrosis was ≤20% of the specimen, and variant histologies (squamous or small cell) were ≤50% (Supplementary Text S1). Clinical and demographic characteristics are described in Supplementary Data File S1.1. Five expert genitourinary
pathologists re-reviewed all of the cases for multiple parameters, including the extent of variant histology (Supplementary Text S1 and Fig. S1.1a).

**Somatic DNA alterations**

The tumours displayed a large number of DNA alterations, slightly fewer than in lung cancer and melanoma, but more than in other adult malignancies studied by TCGA (Fig. 1)\(^9\). On average, there were 302 exonic mutations, 204 segmental alterations in genomic copy number and 22 genomic rearrangements per sample. We analyzed somatic copy number alterations (SCNAs) using both SNP 6.0 arrays and low-pass whole genome sequencing; the two were strongly concordant (Supplementary Methods S6.1 and Supplementary Fig. S6.1). There were 22 significant arm-level copy number changes (Supplementary Data File S6.1.1), and GISTIC (Supplementary Methods S6.2) identified 27 amplified and 30 deleted recurrent focal SCNAs (Supplementary Data Files S6.2.1 and S6.3.1). Focal amplifications involved genes previously reported to be altered in bladder cancer (Fig. 1c, Supplementary Fig. S6.2.1) and some not previously implicated. The latter included *PVRL4, BCL2L1* and *ZNF703*. The most common recurrent focal deletion, seen in 47% of samples, contained *CDKN2A* (9p21.3) and correlated with reduced expression (Fig. 1 and Supplementary Fig. S2.7). Other focal deletions containing <10 genes appeared to target *PDE4D, RB1, FHIT, CREBBP, IKZF2, FOXO1, FAM190A, LRP1B* and *WWOX*.

Whole-exome sequencing of 130 tumours and matched normal samples targeted 186,260 exons in 18,091 genes (mean coverage 100×, with 82% of target bases covered >30×). MuTect\(^10\) identified 39,312 somatic mutations (including 38,012 point mutations and 1,138 indels), yielding mean and median somatic mutation rates of 7.7 and 5.5 per megabase (Mb), respectively (Fig. 1a, Supplementary Table S2.1.1). Thirty-two genes showed statistically significant levels of recurrent somatic mutation (Fig. 1b, Supplementary Table S2.1.2) by analysis using MutSig 1.5\(^11\) (Supplementary Methods S2.2). Three other genes identified by MutSig were not considered further because of low or undetectable expression (Supplementary Fig. S2.1.1). A similar analysis considering only mutations in the COSMIC database\(^2\) identified three more significantly mutated genes (SMGs): *ERBB2, ATM* and *CTNNB1* (Supplementary Table S2.1.3). We validated the mutation findings in three ways: targeted re-sequencing of all SMG mutations, comparison with RNA-Seq data for 123 samples and comparison with whole genome sequence data for 18 samples. Overall, the validation rate was > 99% by a combination of the methods (Supplemental Methods S2.4.)

Nearly half (49%) of the samples had *TP53* mutations (Fig. 1b), which were mutually exclusive in their relationship with amplification (9%) and overexpression (29%) of *MDM2*; hence, TP53 function was inactivated in 76% of samples. Most *RB1* mutations were inactivating, were associated with significantly reduced mRNA level (Supplementary Fig. S2.7) and were mutually exclusive with *CDKN2A* deletions (Supplementary Fig. S2.8 and Table S2.8.1). *FGFR3* mutations (12%) typically affected known kinase-activating sites. *PIK3CA* mutations were relatively common (20%), clustering in the helical domain near E545 (Supplementary Fig. S2.4). Most *TSC1* mutations (8%) were truncating, and six were homozygous (allele fraction > 0.5).
Many of the 32 genes identified in Fig. 1b have not previously been reported as statistically significantly mutated in bladder cancer: MLL2 (27%), CDKN1A* (14%), ERCC2* (12%), STAG2 (11%), RXRA* (9%), ELF3* (8%), NFE2L2 (8%), KLF5* (8%), TXNIP (7%), FOXQ1* (5%), RHOB* (5%), FOXO1 (5%), PAIP1* (5%), BTG2* (5%), ZFP36L1 (5%), RHOA (4%) and CCND3 (4%). The nine genes marked with asterisks have not been reported as SMGs in any other TCGA cancer type or reported in another study as mutated at >3% frequency. CDKN1A (p21CIP1), a cyclin-dependent kinase inhibitor12, had predominantly null or truncating mutations, implying loss of function. Fifteen of 16 mutations in ERCC2, a nucleotide excision repair gene13, were deleterious missense mutations, suggesting dominant negative effects. ERCC2-mutant tumours also had significantly fewer C>G mutations than did ERCC2-wild type tumours (Supplementary Figs. S2.3.1 and S2.3.2), and they trended toward higher overall mutation rate (Supplementary Figure S2.12). Seven of 12 mutations in RXRA (retinoid × nuclear receptor alpha)14 occurred at the same amino acid (five S427F; two S427Y) in the ligand-binding domain. Those seven tumours showed increased expression of genes involved in adipogenesis and lipid metabolism (Supplementary Fig. S2.6 and Data Files S2.6.1–S2.6.3), suggesting that the mutations cause constitutive activation.

Eleven tumours (8%) had deleterious missense mutations in the Neh2 domain of NFE2L2, a transcription factor that regulates the anti-oxidant program in response to oxidative stress15. Those tumours showed dramatically increased expression of genes involved in genotoxic metabolism and the reactive oxygen species (ROS) response (Supplementary Figs. S2.5.1–S2.5.3 and Date File S2.5.2). Furthermore, nine samples had mutations in redox regulator TXNIP16 (5 of them inactivating) and were mutually exclusive of samples with NFE2L2 mutations, providing another mechanism for dysregulation of redox metabolism. Predominant inactivating mutations were seen in STAG2, an X-linked cohesin complex component required for separation of sister chromatids during cell division17 (Supplementary Fig. S2.4).

Unsupervised clustering by non-negative matrix factorization of mutations and focal SCNAs in 125 samples identified three distinct groups (Fig. 1a, Supplementary Fig. S2.1.2). Group A (red), labeled as ‘focally-amplified’, is highly enriched in focal SCNAs in several genes, as well as mutations in MLL2 (Fig. 1; Supplementary Tables S2.1.4 and S2.1.5). Group B (blue), labeled as ‘papillary CDKN2A-deficient FGFR3-mutant’, is enriched in papillary histology. Nearly all Group B samples show loss of CDKN2A, and the majority have one or more alterations in FGFR3. Group C (green), labeled as ‘TP53/cell-cycle-mutant’, shows TP53 mutations in nearly all samples, as well as enrichment with RB1 mutations and amplifications of E2F3 and CCNE1 (Fig. 1, Supplementary Table S2.1.4). Those differences in pattern of mutation suggest the possibility of different oncogenic mechanisms.

Seventy-two per cent of the cancers in this study were from current or past smokers, consistent with extensive epidemiological studies indicating an association between smoking and urothelial cancer risk. In contrast with lung cancer, however, there was no statistically significant association between smoking status and the mutational spectrum, frequency of mutation in any SMG, focal SCNAs or expression subtype (Supplementary Tables S2.9.1 and S2.9.2). Never-smokers did have a slightly higher fraction of C>G mutations than did
current/former smokers (28.5% vs. 23.8%, p = 0.032; Supplementary Figs. 2.3.2 and 2.3.3).

However, unsupervised clustering of promoter CpG island DNA methylation data revealed a
major subgroup (34%) of tumours characterized by cancer-specific DNA hypermethylation
(CIMP) (Supplementary Fig.S7.1). Multivariate regression analysis with age, sex and
tumour stage as covariates identified smoking pack-years as the only significant predictor of
CIMP phenotype, as has also been reported for colorectal cancer\textsuperscript{18}.

Fifty-one per cent of mutations overall were Tp*C->(T/G) (Supplementary Table S2.1.1), a
class of mutation recently reported to be mediated by one of the DNA cytosine deaminases,
APOBEC\textsuperscript{19,20}. APOBEC3B was expressed at high levels in all of the tumours, suggesting a
major role for APOBEC-mediated mutagenesis in bladder carcinogenesis (Supplementary
Figs. S12.1 and S12.2).

Four genes involved in epigenetic regulation were SMGs: MLL2, ARID1A, KDM6A and
EP300 (Fig. 1). Truncating mutations were significantly enriched in each of those genes
(Supplementary Fig. S2.2 and Data Files S2.2.1-2). Three of them had previously been
identified as mutated in urothelial cancers\textsuperscript{4}, but mutation of MLL2, which encodes a histone
H3 lysine 4 (H3K4) methyltransferase, is a novel finding. Several other chromatin-
regulating genes had mutation rates ≥10% but were not statistically significant by MutSig
analysis: MLL3, MLL, CREBBP, CHD7 and SRCAP. Many other epigenetic regulators were
mutated at lower frequency but were also enriched with truncating mutations, suggesting
functional significance (Supplementary Fig. S2.2 and Data Files S2.2.1 and S2.2.2). Non-
silent mutations in chromatin regulatory genes overall were significantly enriched in bladder
cancer in comparison with the entire exome, in contrast with all other epithelial cancers
studied to date in the TCGA project (Supplementary Table S2.10). Mutations in MLL2 and
KDM6A (the latter encoding a histone H3 lysine 27 (H3K27) demethylase) were mutually
exclusive (Supplementary Fig. S2.8 and Table S2.8.1), suggesting that mutations in the two
genes have redundant downstream effects on carcinogenesis or that the combined loss is
synthetically lethal.

**Chromosomal rearrangements and viral integration**

To identify structural variations and pathogen sequences, we used low-pass, paired-end,
whole-genome sequencing (WGS; 6-8x coverage) of 114 tumours and RNA sequencing of
all tumours. We detected 2,529 structural aberrations, including 1,153 that involve gene-
gene fusions. Among the translocations, 379 were inter-chromosomal, 237 were intra-
chromosomal, 274 were the result of inversions and 263 resulted from deletions
(Supplementary Table S3.1). We found several recurrent translocations of likely pathogenic
significance, including an intra-chromosomal translocation on chromosome 4 involving
FGFR3 and TACC3 (n=3). The breakpoints were in intron 16 (2 cases) or exon 17 (1 case)
of FGFR3 and intron 10 of TACC3 (confirmed by DNA sequencing and RNA-seq). All
three lead to fusion mRNA products whose predicted proteins include the N-terminal 758
amino acids of FGFR3 fused with the C-terminal 191 amino acids of TACC3 (Fig. 2a).

Based on the structure of the FGFR3-TACC3 fusion protein, we predict that it can auto-
dimerize, leading to constitutive activation of the kinase domain of FGFR3. FGFR3-TACC3
fusion, which was recently described in both glioblastoma\textsuperscript{21} and bladder cancer\textsuperscript{7,8}.
represents a promising therapeutic target. The ERBB2 gene was also involved in translocations in four tumours, all with different fusion partners and all confirmed by DNA sequencing, RNA-Seq or both. In one case, exons 4 to 29 of ERBB2 were fused to the promoter plus exon 1 of DIP2B, and the fusion product was amplified (Fig. 2b). Two other fusion products resulted in novel mRNA products whose biological significance is not known.

We identified viral DNAs in 7 of 122 tumours (6%), and viral transcripts in 5 of 122 (4%). Three tumours expressed cytomegalovirus (CMV) transcripts (encoding RL5A, RNA2.7, RL9A, RNA1.2, UL5 and UL22A), one expressed BK polyoma virus and one expressed human papilloma virus 16 (HPV16). HPV16 and human herpes virus 6B DNA were each identified in one other sample but without expression. None of the tumours expressing CMV showed evidence of CMV integration into the host genome, suggesting the presence of a stable episome. In the BK-positive tumour, two BK genes were integrated into GRB14, a signaling adapter protein for receptor tyrosine kinases. In the HPV 16-expressing case, the virus integrated into BCL2L1, an apoptosis-regulating gene (Fig. 2c). In that tumour, BCL2L1 was amplified (~6×) and overexpressed (~10× median; > 2× any of the other samples). Overall, those findings suggest that viral infection may play a role in the development of a small percentage of urothelial carcinomas.

**mRNA, microRNA and protein expression**

Analysis of RNA-seq data from 129 tumours identified four clusters (clusters I-IV) (Fig. 3 and Supplementary Fig. S4.1). Cluster I (‘papillary-like’) is enriched in tumours with papillary morphology (p=0.0002), FGFR3 mutations (p=0.0007, q=0.02), FGFR3 copy number gain (p=0.04, q=0.1) and elevated FGFR3 expression (p<0.0001) (Fig. 3a). It includes all three samples with FGFR3-TACC3 fusions. Cluster I samples also show significantly lower expression of miR-99a and miR-100, microRNAs that down-regulate FGFR3 expression (p=0.0002, Figs. 3a S5.3)22. They also show lower expression of miR-145 and miR-125b, which have been reported as frequently downregulated in bladder cancer23. Tumours with FGFR3 alterations, and perhaps other tumours that share the Cluster I expression profile, may respond to inhibitors of FGFR or its downstream targets.

Reverse-phase protein array (RPPA) data indicate that clusters I and II express high HER2 (ERBB2) levels and an elevated estrogen receptor beta (ESR1) signaling signature, indicating potential targets for hormone therapies such as tamoxifen or raloxifene (Fig. 3d). In fact, the HER2 protein levels in a subset of the tumours are comparable to those found in TCGA HER2-positive breast cancers21.

For comparison, we asked whether any of the four clusters show gene signatures similar to those identified in any other tumour type(s) among the first 11 analyzed by TCGA. We found that the signature of bladder cancer cluster III (‘basal/squamous-like’) is similar to that of basal-like breast cancers, as well as squamous cell cancers of the head and neck and lung (Supplementary Fig. S4.2)24,25. All four of those cancer types express characteristic epithelial lineage genes, including KRT14, KRT5, KRT6 and EGFR. Basal-like subtype26 and squamous cell subtype27 of urothelial carcinoma, have been independently reported. Many of the samples in bladder cluster III express cytokeratins (i.e. KRT14 and KRT5) that
were recently reported to mark stem/progenitor cells. Some of those samples also show a level of variant squamous histology (Fig. 3b). Bladder clusters I and II show features similar to those of Luminal A breast cancer, with high mRNA and protein expression of luminal breast differentiation markers, including GATA3 and FOXA1 (Fig. 3c). Markers of urothelial differentiation such as the uroplakins are also highly expressed in clusters I and II, as are the epithelial marker E-cadherin and members of the miR-200 family of microRNAs (which target multiple regulators of epithelial-mesenchymal transition) (Fig. 3c). Taken together, those observations suggest that, despite their diverse tissue origins, some bladder, breast, head and neck and lung cancers share common pathways of tumour development.

To determine if the expression-based clusters could be seen in other datasets, we used the muscle-invasive bladder cancer samples from Sjodahl, et al., hierarchically clustering them with the genes used in our analysis. From the sample dendrogram, we identified four groups (Supplemental Figure S4.3a). The four groups identified in the Sjodahl data set correlated well with the four clusters identified in our TCGA data. (Supplemental Figure S4.3b).

When we analyzed the RNA-seq data for transcript splice variation using SpliceSeq (Supplemental Material S11), one finding of interest was an average of 3% PKM1 and 97% PKM2 transcripts in the tumour samples. The PKM2 isoform of pyruvate kinase is the principal driver of a shift to aerobic glycolysis in tumours (the Warburg effect). Therefore, urothelial bladder cancers (and other cancer types) may prove sensitive to inhibition of glycolysis or related metabolic pathways.

Pathway analysis and therapeutic targeting

Integrated analysis of the mutation and copy-number data revealed three main pathways as frequently dysregulated in bladder cancer: cell cycle regulation (altered in 93% of cases); kinase and PI3K signaling (72%); chromatin remodeling, including mutations/SCNAs in histone modifying genes (89%); and components of the SWI/SNF nucleosome remodeling complex (64%) (Fig. 4a). To complement those results for well-defined pathways, we applied network analysis methods to examine other possible interactions between genes and pathways (Fig. 4b). In particular, we used the TieDIE algorithm to search for causal regulatory interactions within the PARADIGM network, which connects mutated genes to active transcriptional hubs. The analysis identified a sub-network linking mutated histone-modifying genes to a large array of activated transcription factors, suggesting potential far-reaching effects of histone modification on other pathways (Supplementary Fig. S8.2.1), converging on MYC/MAX regulation. Both MYC and MAX showed similar levels of pathway activity, independent of mutations in chromatin genes, suggesting that mutations in histone-modifying genes provide just one mechanism for disruption of the MYC/MAX hub. In contrast, tumours with chromatin-related mutations showed differential activity of transcription factors FOXA2 and SP1, implicating de-differentiation processes as a result of the mutations. Our network analysis also identified HSP90AA1 as a critical signaling hub, suggesting that inhibitors of HSP90 may have therapeutic value in urothelial carcinoma.

Although the linkages between mutations and transcriptional changes were statistically significant in terms of their proximity in the network (as determined by permutation tests;
see Supplementary Fig. S8.2), further studies will be needed to assess the biological relevance of the findings.

Integrated analysis also identified mutations, copy number alterations or RNA expression changes affecting the PI3-kinase/AKT/mTOR pathway in 42% of the tumours (Fig. 5a). Included were activating point mutations in PIK3CA (17%; potentially responsive to PI3K inhibitors), mutation or deletion of TSC1 or TSC2 (9%; potentially responsive to mTOR inhibitors) and overexpression of AKT3 (10%; potentially responsive to AKT inhibitors). We also observed mutations, genomic amplifications or gene fusions that affect the RTK/RAS pathway in 44% of the tumours (Fig. 5b). Included were events that can activate FGFR3 (17%; potentially responsive to FGFR inhibitors or antibodies), amplification of EGFR (9%; potentially responsive to EGFR antibodies or inhibitors), mutations of ERBB3 (6%; potentially sensitive to ERBB kinase inhibitors) and mutation or amplification of ERBB2 (9%; potentially sensitive to ERBB2 kinase inhibitors or antibodies). ERBB3 mutations in bladder cancer have been noted previously but statistically significant mutation of ERBB2 in bladder cancer has not been reported. Both genes are potential therapeutic targets in other diseases (Fig. 5c). Interestingly, ERBB2 alterations were approximately as frequent in this study as in TCGA breast cancers, but with fewer amplifications and more mutations (Fig. 5d).

Discussion

This integrated study of 131 invasive urothelial bladder carcinomas provides numerous novel insights into disease biology and delineates multiple potential opportunities for therapeutic intervention. Treatment for muscle-invasive bladder cancer has not advanced beyond cisplatin-based combination chemotherapy and surgery in the past 30 years, and no new drugs for the disease have been approved in that time. Median survival for patients with recurrent or metastatic bladder cancer remains 14-15 months with cisplatin-based chemotherapy, and there is no widely recognized second-line therapy. With the exception of a single case report, there is also no known benefit from treatment with newer, targeted agents. Several of the genomic alterations identified in this study, particularly those involving the PI3-kinase/AKT/mTOR, CDKN2A/CDK4/CCND1 and RTK/RAS pathways, as well as ERBB2 (Her-2), ERBB3 and FGFR3, are amenable in principle to therapeutic targeting. Clinical trials based on patients with relevant druggable genomic alterations are warranted.

FGFR3 mutation is a common feature of low-grade non-invasive papillary urothelial bladder cancer, but it occurs at a much lower frequency in high-grade invasive bladder cancer. The cluster analysis in Figure 3 highlights multiple mechanisms of FGFR3 activation, and its strong association with papillary morphology. The data presented here suggest a subset of muscle-invasive cancers that can potentially be targeted through FGFR3. Similarly, ERBB2 amplification may be targetable by strategies used in breast cancer, by small-molecule tyrosine kinase inhibitors or by novel immunotherapeutic approaches (NCT01353222). The data here provide further support for several ongoing ERBB2-targeted trials in bladder cancer and further define the subpopulation of cancers suited to that approach. Finally, cluster III of the integrated expression profiling analysis reveals the existence of a urothelial
carcinoma subtype with high cancer stem cell content (including KRT14/5), perhaps providing another avenue for therapeutic targeting.

The alterations identified in epigenetic pathways also suggest new possibilities for bladder cancer treatment. Ninety-nine (76%) of the tumours analyzed here had an inactivating mutation in one or more of the chromatin regulatory genes, and 53 (41%) had at least two such mutations. Overall, the bladder cancers showed a mutational spectrum highly enriched with mutations in chromatin regulatory genes (Supplementary Table S2.10). Further, integrated network analyses revealed a profound impact of those mutations on the activity levels of various transcription factors and pathways implicated in cancer. Recent development of drugs that bind competitively to acetyl-lysine recognition motifs (i.e., bromodomains) might prove useful for treatment of the subset of bladder tumours that have abnormalities in chromatin-modifying enzymes\textsuperscript{39}. Our findings suggest that bladder cancer is a prime candidate for further exploration of that approach to therapy.

Methods Summary

Tumour and normal samples were obtained with institutional review board-approved consent and processed using a modified AllPrep kit (Qiagen) to obtain purified DNA and RNA. Quality-control analyses revealed only modest batch effects (Supplementary Text S13.1). The tumours were profiled using Affymetrix SNP 6.0 microarrays for SCNAs, low-pass WGS (HiSeq) for SCNAs and translocations, RNA-seq (HiSeq) for mRNA and miRNA expression, Illumina Infinium (HumanMethylation450) arrays for DNA methylation, HiSeq for exome sequencing and RPPA for protein expression and phosphorylation. Statistical analysis and biological interpretation of the data were spearheaded by the TCGA Genome Data Analysis Centers. Sequence files are in CGHub (https://cghub.ucsc.edu/). All other molecular, clinical and pathological data are available through the TCGA Data Portal (https://tcga-data.nci.nih.gov/tcga/). Data matrices, molecular analysis results and supporting information are at http://tcga-data.nci.nih.gov/docs/publications/bladder_2013/. The data can be explored through a compendium of Next-Generation Clustered Heat Maps (http://bioinformatics.mdanderson.org/main/TCGA/Supplements/NGCHM-BLCA), the cBio Cancer Genomics Portal (http://cbióportal.org), PARADIGM (http://sysbio.soe.ucsc.edu/paradigm/tutorial/), SpliceSeq (http://bioinformatics.mdanderson.org/main/SpliceSeq:Overview), MBatch batch effects assessor (http://bioinformatics.mdanderson.org/tcgbatcheffects) and Regulome Explorer (http://explorer.cancerregulome.org/). Also see Supplementary Materials.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. The genomic landscape of bladder cancer

a. Mutation rate and type, histologic subtype, smoking status, gender, tumour stage, and cluster type. 
b. Genes with statistically significant levels of mutation (MutSig, FDR < 0.1) and mutation types. 
c. Deletions and amplifications for genomic regions with statistically significant focal copy number changes (GISTIC2.0). CN refers to absolute copy number. 
   Note that two amplification peaks (*) contain several genes, any of which could be the target, as opposed to the single gene listed here.
d. RNA expression level expressed as fold change from the median (of all samples). RPKM values are shown for selected genes subject to mutation and/or focal copy number change. Tumour samples were grouped into three clusters (red, blue, and green) using consensus NMF clustering (see the main text and Supplementary Figure 2.1.2). Three samples with no copy number data and two samples with no mutations in the genes were not used in the clustering and are shown in gray.
Figure 2. Structural rearrangements and viral integration

a. FGFR3-TACC3 fusion in sample TCGA-CF-A3MH showing the breakpoints in the two genes, the breakpoint junction sequences and the predicted fusion protein.

b. Rearrangement involving DIP2B and ERBB2 in TCGA-DK-A2I6. The ERBB2 gene has swapped its promoter with that of DIP2B, resulting in over-expression of ERBB2.

c. Insertion of human papilloma virus 16 (HPV16) into the BCL2L1 gene on chromosome 20 in TCGA-GC-A3I6. The region of BCL2L1 into which the virus has integrated and the integration junction sequence are shown.
Figure 3. Expression characteristics of bladder cancer
Integrated analysis of mRNA, miRNA and protein data led to identification of distinct subsets of urothelial carcinoma. Data for mRNA, miRNA and protein were z-normalized, and samples were organized in the horizontal direction by mRNA clustering. a, Papillary histology, FGFR3 alterations, FGFR3 expression and reduced FGFR3-related miRNA expression are enriched in cluster I. b, Expression of epithelial lineage genes and stem/progenitor cytokeratins are generally high in cluster III, some of which express variant squamous histology. c, Luminal breast and urothelial differentiation factors are enriched in clusters I and II. d, ERBB2 mutation and estrogen receptor beta (ESR2) expression are enriched in clusters I and II.

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Figure 4. Altered pathways and networks in bladder cancer

a. Somatic mutations and copy number alterations (CNA) in components of the p53/Rb pathway, RTK/RAS/PI3K pathway, histone modification system and SWI/SNF complex. Red, activating genetic alterations; blue, inactivating genetic alterations. Percentages shown denote activation or inactivation of at least one allele. 
b. The network connecting mutated histone-modifying genes to transcription factors with differential activity (methodology and larger implicated network in Supplementary Fig. 8.2.1). Each gene is depicted as a multi-ring circle with various levels of data, plotted such that each ‘spoke’ in the ring represents a single patient sample (same sample ordering for all genes). ‘PARADIGM’ ring: bioinformatically inferred levels of gene activity (red, higher activity); ‘Transcriptional Activity’: mean mRNA levels of all of the targets of each transcription factor; ‘Expression’: mRNA levels relative to normal (red, high); ‘Mutation in gene’: somatic mutation; ‘Mutation in histone modifier genes’: somatic mutation in at least one such gene; ‘IPL anti-correlation’: genes with PARADIGM Integrated Pathway Levels (IPLs) inversely correlated with histone-gene mutation status. Gene-gene relationships inferred using public resources.
Figure 5. Potential targets in bladder cancer

a. Alterations in the PI3K/AKT/mTOR pathway are mutually exclusive. Tumour samples are shown in columns, genes in rows. Only samples with at least one alteration are shown. AKT3 shows elevated expression in 10% of samples, independent of copy-number (right panel).

b. Receptor tyrosine kinases are altered, by any of several different mechanisms (amplification, mutation, and fusion), in 45% of samples. Only mutations that are recurrent in this data set or previously reported in COSMIC are shown.

c. Recurrent mutations in ERBB2 and ERBB3. The mutations shown in black are either recurrent in the TCGA data set or are reported in COSMIC. Green: Receptor L domain; red: furin-like cysteine-rich region; blue: growth factor receptor domain IV; yellow: tyrosine kinase domain.

d. ERBB2 amplifications and recurrent mutations in other projects profiled by TCGA. Missense mutations were counted in the following positions: G309, S310, L313, R678, T733, L755, V777, D769, V842, T862, R896, M916I. In-frame insertions were counted between amino acids 774 and 776. Only tumour types with an alteration frequency ≥ 2% are shown.