Multi-omics Perspective on the Tumor Microenvironment based on PD-L1 and CD8 T-Cell Infiltration in Urothelial Cancer

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Abstract

Objectives: We carried out an integrated analysis based on multiple-dimensional types of data from cohorts of bladder cancer patients to identify multi-omics perspective (genomics and transcriptomics) on the tumor microenvironment on the bases of the programmed cell death 1 ligand (PD-L1) and CD8 T-cell infiltration in urothelial carcinoma.

Methods: Multiple-dimensional types of data, including clinical, genomic and transcriptomic data of 408 bladder cancer patients were retrieved from the Cancer Genome Atlas database. Based on the median values of PD-L1 and CD8A, the tumor samples were grouped into four tumor microenvironment immune types (TMIT). The RNA sequencing profiles, somatic mutation and PD-L1 amplification data of bladder cancer were analyzed by different TMITs.

Results: Our research demonstrated that 36.8% of the evaluated bladder cancer belonged to TMIT I (high PD-L1/high CD8A). TMIT subtypes were not significantly associated with overall survival or disease free survival in urothelial cancer. TMIT I facilitates CD8+ T-cell infiltration and activates T-effector and interferon gamma (IFN-γ) associated gene signature. The number of somatic mutations, cytolytic activity, IFN-γ mRNA expression and TIGIT mRNA expression in TMIT I was remarkably higher than those in other TMIT groups. Our results showed a high rate of C>T transversion and a high rate of transition/transversion (Ti/Tv) in TMIT I bladder tumors. The RB1 mutation was significantly associated with TMIT I bladder cancer and be significantly co-occurring with the TP53 mutation. However, FGFR3 mutation and TP53 mutation were mutually exclusive in TMIT II bladder tumors. More importantly, different amino acid changes by FGFR3/RB1 mutations were also found between TMIT I and TMIT II bladder cancer, such as amino acid changes in "Immunoglobulin I-set domain (260-356)" and "Protein tyrosine kinase (472-748)". We also detected 9 genes as significantly cancer-associated genes in TMIT I bladder cancer, of which, RAD51C has been reported to play an important role in DNA damage responses. Further analysis concentrated on the potential molecular mechanism found that TMIT I was significantly associated with anti-tumor immune-related signaling pathway, and kataegis was present on chromosome 21 in TMIT I bladder tumors.

Conclusions: The classification of bladder cancer into four TMITs on the bases of the PD-L1 expression and the CD8+ CTLs statuses is an appropriate approach for bladder tumor immunotherapy. TMIT I (high PD-L1/high CD8A) is significantly correlated with more somatic mutation burden, and facilitates CD8+ T-cell infiltration and activates T-effector and IFN-γ associated gene signature. Alteration landscape for somatic variants was different between TMIT I and TMIT II (low PD-L1/low CD8A).

Key words: Bladder cancer, PD-L1, Immunotherapy, TMIT, TIGIT, kataegis

Introduction

Recent attempts of immune checkpoint inhibitors that target the programmed cell death 1 (PD-1) and PD-1 ligand (PD-L1) have revealed robust antitumor activity with a favorable safety profile in
the treatment of recurrent advanced urothelial carcinoma [1, 2]. However, the response rates are relatively low, and there are still no reliable predictive biomarkers of clinical benefits of anti–PD-1/PD-L1 therapies in bladder carcinoma to date.

Encouragingly, recent studies have prominently highlighted that PD-L1 expression [3], tumor mutational load [4], and the intensity of CD8+ T-cell infiltrates [5] could be potential biomarkers of clinical benefits of anti–PD-1/PD-L1 therapies. Furthermore, the classification of carcinoma into four different types on the bases of the CD8+ cytotoxic T cells (CTL) statuses and the expression of PD-L1 has been proposed recently [6]. Consistent with previous observations, cancers with the presence of CD8+ CTLs and high level of PD-L1 expression belong to tumor microenvironment immune type (TMIT) I, which was proved to be the potential predictive biomarker of clinical response to PD-L1/PD-1 immunotherapy [7]. However, this novel notion has not been verified by a bladder cancer genomics profile.

Here, we carry out an integrated analysis on the bases of multiple-dimensional types of data, including clinical, genomic and transcriptomic data from cohorts of bladder cancer patients to identify multi-omics perspective on the tumor microenvironment on the bases of CD8 T-Cell infiltration and PD-L1 expression in urothelial carcinoma.

Materials and Methods

Demographic, clinical and pathological data

The Cancer Genome Atlas (TCGA) cohort was retrieved from the online data repository. A total of 408 bladder cancer samples were recruited in TCGA series with gene mutation data, mRNA expression profiles, and gene copy number alteration data. Key variables, including clinical and demographic information, were provided in Table 1.

Somatic variants data, mRNA expression profiling, and gene copy number alteration analysis

Analysis of RNA sequencing data from 408 bladder cancers was retrieved from the TCGA database. Experimental procedures, including RNA sequencing, quality control, mRNA library reading and further quantification of gene expression have been reported previously [8]. The cytolytic activity was defined as the geometric mean value of granzyme and perforin1 in each sample [9]. Mutation annotation format (MAF) of the 408 bladder cancers, which was used to store somatic variants detected, were also downloaded from the TCGA database. We used the package of “maftools” to summarize, analyze, annotate and visualize MAF files in an efficient manner [10]. Mutation spectrum for each tumor was evaluated as the percentage of six possible single nucleotide changes among single nucleotide substitutions as previously reported [11]. The predicted neoantigen number was referenced in a previous report by Chan-Young Ock [6]. Gene copy number alteration (CNA) analysis was based on CNA data from the TCGA database. Affymetrix SNP6.0 arrays were used to assess somatic CNAs and putative copy-number called in 408 cases determined using GISTIC 2.0. Values as previously described [8].

Table 1. Demographic, clinical and pathological information of TCGA cohort.

| TCGA Cohort          | No. of patients | Age  | Gender | Stage | TMIT types | FGFR3 | mRNA burden | CD8A | PD-L1 | IFN-γ | Neoantigen number |
|----------------------|-----------------|------|--------|-------|------------|-------|-------------|------|-------|-------|------------------|
|                      |                 | Mean | n(%)   | n(%)  | n(%)       | n(%)  | n(%)        |      |       |       |                  |
|                      |                 | 75.0 | 150(36.8) | 107(26.2) | 337(82.6) | 356.9±384.7 | 3.1±5.8 | 3.1±5.1 | 0.6±1.5 | 6.6±1.8 | 45.8±42.8        |
|                      |                 | 40±98| 130(31.9) | 140(34.3) | 134(32.8) | 150(36.8) | 150(36.8) | 54(13.2) | 54(13.2) | 54(13.2) | 54(13.2)         |
|                      |                 | 301(73.8) | 107(26.2) | 2(0.5) | 2(0.5) | 107(26.2) | 301(73.8) | 107(26.2) | 2(0.5) | 2(0.5) | 2(0.5)          |
|                      |                 | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) |
|                      |                 | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) |
|                      |                 | 360(14.7) | 360(14.7) | 360(14.7) | 360(14.7) | 360(14.7) | 360(14.7) | 360(14.7) | 360(14.7) | 360(14.7) | 360(14.7) |
|                      |                 | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  |
|                      |                 | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  | 60(14.7)  |
|                      |                 | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) | 348(85.3) |
|                      |                 | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) |
|                      |                 | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) |
|                      |                 | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) | 337(82.6) |

The cytolytic activity of each sample was calculated using the value of the geometric mean of granzyme and perforin1. The predicted neoantigen number was referenced in a previous report by Chan-Young Ock. TCGA, the Cancer Genome Atlas; TMIT, tumor microenvironment immune type; x±σ, mean ± standard deviation.

All of the 408 bladder cancer samples belonged to four TMITs as follows: type I, with expressions level of PD-L1 and CD8A higher than their median values, respectively; type II, with expressions level of PD-L1 and CD8A lower than their median values, respectively; type III, with higher PD-L1 expression and lower CD8A expression in contrast to their median values, respectively; and type IV, with lower PD-L1 expression and higher CD8A expression in contrast to their median values, respectively [7].
Gene set enrichment analysis

We also performed “Gene Set Enrichment Analysis (GSEA)” based on the molecular signatures database C5 BP: GO biological process collection [12], to associate the signature of genes with the TMIT I bladder tumors. False discovery rate (FDR) was calculated to effectively control false positives proportion. The normalized enrichment score (NES) was regarded as the primary statistic for examining GSEA enrichment results. The significance threshold was set at FDR < 0.0001.

Statistical analysis

In our study, the categorical variable was analyzed as a percentage and compared by a Pearson’s Chi-square test. The continuous variable was analyzed using either a two-tailed Student’s t-test or a one-way analysis of variance. The significant relationship between two continuous variables, such as the cytolytic activity, the expression level of interferon gamma (IFN-γ) mRNA and CD8A mRNA, was calculated by linear regression analysis. We also performed logistic regression analyses to determine the potential statistically significant relationship between genomic characteristics and the prediction of TMIT I. Statistical Package for Social Sciences 24.0 software (SPSS Inc., Chicago, IL, USA) was performed for statistical analysis. P values less than 0.05 were considered significant.

Results

A total number of 408 bladder cancer samples, included in the TCGA cohort, were grouped into four TMITs based on the expression levels of CD8A and PD-L1. As shown in Fig. 1a, 36.8% of the evaluated bladder tumors belonged to TMIT I, with a high proportion of CD8+ CTLs and high expression level of PD-L1. The proportions of TMIT II, III, and IV were 36.8%, 13.2%, and 13.2%, respectively. TIMT subtypes were not significantly associated with overall survival (OS) or disease free survival (DFS) in urothelial cancer (Fig. S1). The detailed clinical, pathological and molecular characteristics for the four TMIT types of the tumor samples were shown in Table 2.

![Fig. 1. Landscape of TMIT, mutational signatures and mutations. (a) Scatter plot for the distribution of TMITs. (b) Comparison of somatic mutation burden between different TMITs. (c) Comparison of the proportion of TMIT I according to somatic mutation burden. (d) Boxplot for classification of somatic variants in 150 TMIT I bladder tumors. (e) Alteration landscape for 150 TMIT I bladder tumors and 150 TMIT II bladder tumors. TMIT, tumor microenvironment immune type; Ti, transition; Tv, transversion; ***, P < 0.001; **, P < 0.01; *, P < 0.05.](http://www.jcancer.org)
been reported to be associated with the immunotherapy to tumors [9, 13] . Here, we predicted neoantigen in bladder cancer. Notably, a correlation between somatic mutation burden and performed a linear regression analysis to explore the correlation between somatic mutation burden and the number of neoantigens was found (R²=0.767, P<0.0001, Fig. S2). More importantly, TMIT I group had a remarkable higher somatic mutation burden compared with the higher proportion of TMIT I, compared with wild-type group (13.3% vs.40.8%; P < 0.001; Fig. S3a), TMIT II tumors (412.2 vs.297.0; P < 0.01, Fig. 1b). Bladder tumors with more somatic mutation burden (higher than the median value) were accompanied with the higher proportion of TMIT I, compared with those with fewer mutations (44.6% vs.28.9%; P < 0.01; Fig. 1c).

**High somatic mutation burden were correlated with TMIT I**

The total mutation count and neoantigen have been reported to be associated with the immunotherapy to tumors [9, 13]. Here, we performed a linear regression analysis to explore the correlation between somatic mutation burden and predicted neoantigen in bladder cancer. Notably, a statistically significant correlation between the somatic mutation burden and the number of neoantigens was found (R²=0.767, P<0.0001, Fig. S2). More importantly, TMIT I group had a remarkable higher somatic mutation burden compared with the TMIT II tumors (412.2 vs.297.0; P < 0.01, Fig. 1b). Bladder tumors with more somatic mutation burden (higher than the median value) were accompanied with the higher proportion of TMIT I, compared with those with fewer mutations (44.6% vs.28.9%; P < 0.01; Fig. 1c).

**Alteration landscape for somatic variants was different between TMIT I (high PD-L1/high CD8<sub>8</sub>) and TMIT II (low PD-L1/low CD8<sub>8</sub>)**

We sought to explore whether TMIT I was associated with specific tumor mutation spectrum. Our results showed a high rate of C>T transversion and a high rate of transition/transversion (T/T vs.) in TMIT I bladder tumors (Fig. 1d). Alteration landscape for 150 TMIT I bladder tumors and 150 TMIT II bladder tumors were shown in Fig. 1e. Seven genes were mutated in >25% of samples in TMIT I: TTN (62%), TP53 (57%), KMT2D (29%), MUC16 (29%), RB1 (27%), HMCN1 (25%), and PIK3CA (25%). While six genes were mutated in >25% of samples in TMIT II: TTN (43%), TP53 (41%), KDM6A (31%), KMT2D (29%), FGFR3 (25%) and ARID1A (25%). Specifically, a higher rate of FGFR3 mutation in TMIT II bladder tumors and a higher rate of RB1 mutation in TMIT I bladder tumors were found.

Interestingly, the vast majority of tumors with FGFR3 mutation were classified as TMIT II (low PD-L1/low CD8<sub>8</sub>) (Fig. 2a). Similarly, the proportion of TMIT I in FGFR3 mutated tumors was statistically significantly lower than that in FGFR3 wild-type group (13.3% vs.40.8%; P < 0.001; Fig. S3a), consistent with the distribution of FGFR3 mutations in the scatter plot. However, the high proportion of tumors with RB1 mutation were grouped into TMIT I (Fig. 2c), and the proportion of TMIT I in RB1 mutated tumors was statistically significantly higher than that in RB1 wild-type group (56.3% vs. 32.6%; P < 0.001; Fig. S3b).

In consideration of the actual discovery that FGFR3/RB1 mutations were significantly correlated different between TMIT I and TMIT II bladder tumors, our study next to explore whether there were different amino acid changes by FGFR3/RB1 mutations between TMIT I and TMIT II bladder cancer. Our labelling points revealed that there were different amino acid changes in “Immunoglobulin
I-set domain (260-356) “and “Protein tyrosine kinase (472-748)” by FGFR3 mutation between TMIT I and TMIT II bladder tumors (Fig. 2b, Table 3). We also found different amino acid changes in “Retinoblastoma-associated protein A domain (373-573)”, “Retinoblastoma-associated protein B domain (645 - 766)” and “Rb C-terminal domain (768 - 927)” by RB1 mutation between these two subgroups (Fig. 2d, Table 3).

Many disease-causing genes in cancer are co-occurring or show strong exclusiveness in their mutation pattern. We next explored the potential different somatic interactions between TMIT I and TMIT II bladder tumors (Fig. 3a). The results demonstrated that in TMIT I bladder tumors, RB1 mutation and TP53 mutation were found to be significantly co-occurring (P < 0.05). While in TMIT II bladder tumors, FGFR3 mutation was proved to be significantly associated with STAG2 mutation and KDM6A mutation (P < 0.05). More importantly, FGFR3 mutation and TP53 mutation were mutually exclusive in TMIT II bladder tumors (P < 0.05).

Most of the variants in cancer-causing genes are enriched at few specific loci. To detect such cancer-associated genes, we performed a function called “oncodrive”, based on oncodriveCLUST algorithm originally implemented in Python framework [10, 14]. On the TMIT I cohort, oncodrive was able to detect 9 genes as significantly disease-associated genes, including RAD51C and PLK3. While on the TMIT II cohort, we detect 6 genes as significantly disease-associated genes, including FGFR3 and KRAS (Fig. 3b, Table 3).

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Table 3. Different amino acid changes and related cancer-driver genes detected in TMIT I and TMIT II bladder tumors.

| TMIT I | Mutant gene (%) | Different amino acid changes | Related cancer-driver genes |
|--------|----------------|-----------------------------|-----------------------------|
| I      | RB1 (26.7%)    | Retinoblastoma-associated protein A domain (373-573) | ZNF878, RCPD8, EEF1B2, KRTAP1-5, XRRA, GNA13, RAD51C, KRTAP4-11, PLK3 |
|        |                | Retinoblastoma-associated protein B domain (645 - 766) |                                          |
|        |                | Rb C-terminal domain (768 - 927) |                                          |
| II     | FGFR3 (25.3%)  | Immunoglobulin I-set domain (260-356) | ZNF814, FGFR3, POTEM, KRAS, RCPD8, RHOA |
|        |                | Protein tyrosine kinase (472-748) |                                          |
Fig. 3. Comparison of somatic interactions, cancer driver genes and genomic loci with localized hyper-mutations between TMIT I and TMIT II bladder tumors. (a) Heatmap of mutually exclusive or co-occurring set of genes in the mutation pattern of TMIT I and TMIT II bladder tumors. (b) Detecting cancer driver genes based on positional clustering in TMIT I and TMIT II bladder tumors. Each dot represents a gene and size of the dot represents number clusters (mentioned inside square brackets) within which, a fraction (X-axis) of total variants are accumulated. (c) Rainfall plots for the genomic loci with localized hyper-mutations by inter variant distance on a linear genomic scale. Each dot represented a single nucleotide variants (SNV) and were color coded according to six substitution classes. Arrowheads indicated clusters of hyper mutated genomic regions called as “kataegis”. TMIT, tumor microenvironment immune type.

Table 4. Hyper mutated genomic regions detected in TMIT I and TMIT II bladder tumors.

| TMIT | Chromosome | Start Position | End Position | nMuts | Avg inter-mutation dist | Size | C > A | C > G | C > T | T > C | T > A | T > G |
|------|------------|----------------|--------------|-------|-------------------------|------|-------|-------|-------|-------|-------|-------|
| I    | 21         | 16339706       | 43867289     | 15    | 1966256                 | 27527583 | NA    | 7     | 8     | NA    | NA    | NA    |
| II   | 1          | 150689482      | 152283878    | 25    | 66424.83                | 1594196 | 5     | 13    | 6     | 1     | NA    | NA    |
|      | 15         | 41859720       | 42442033     | 6     | 116462.6                | 582313  | NA    | 4     | 2     | NA    | NA    | NA    |

nMuts, number of mutations; Avg inter-mutation dist, average inter-mutation distance
Cancer genomes have been reported to be characterized by hypermutated genomic regions called as “kataegis” [15]. We next explored the clusters of mutations in kataegis in both TMIT I and TMIT II cohorts. Rainfall plots for the genomic loci with localized hyper-mutations indicated that kataegis was present on chromosome 21 in TMIT I bladder tumors. However, kataegis was present on chromosomes 1 and 15 in TMIT II bladder tumors (Fig. 3c, Table 4).

**TMIT I facilitates CD8+ T-cell infiltration and activates T-effector and IFN-γ associated gene signature**

Then we tried to explore the potential relationship between TMIT I and CD8+ T-cell infiltration and IFN-γ associated gene signatures of the bladder cancer. And the results revealed that high proportions of tumors with high IFN-γ mRNA expression or high cytolytic activity were grouped into TMIT I, compared with other TMIT groups (Fig. 4a and Fig. 4i). Linear regression analysis found that IFN-γ mRNA expression was statistically significantly correlated with the CD8A mRNA expression (R²=0.598, P<0.0001, Fig. 4d). More importantly, both of the IFN-γ mRNA expression and the cytolytic activity were strikingly higher in TMIT I tumors than other TMIT groups (P < 0.001; Fig. 4b and Fig. 4g).

To verify the correction between IFN-γ mRNA expression and TMIT, the proportion of TMIT I was compared based on IFN-γ mRNA expression. We found that bladder tumors with a higher expression of IFN-γ mRNA had a remarkably higher proportion of TMIT I (68.1% vs 5.4%; P < 0.001; Fig. 4c). Similarly, bladder tumors with a higher cytolytic activity than the median value had a notably higher proportion of TMIT I, compared with those with lower cytolytic activity (67.6% vs 5.9%; P < 0.001; Fig. 4h).

Next, we sought to find the relationship between PD-L1 amplification and TMIT by CNA analysis
based on the TCGA database. The proportion of PD-L1 amplification in bladder cancers was 21.1% (86/408). The proportion of TMIT I in the tumors with PD-L1 amplification was statistically higher than that in the control group (66.3% vs. 29.2%; P < 0.001; Fig. 4e). Remarkably, scatter plot for the distribution of PD-L1 amplification demonstrated that almost all of the bladder cancers with PD-L1 amplification were classified as TMIT I, compared with other TMIT groups (Fig. 4f), which was consistent with observations in Fig. 4e.

We next explored the potential relationship of TIGIT mRNA expression and TMIT I in bladder cancer. Scatter plot for the distribution of TIGIT mRNA expression revealed that a high proportion of tumors with high TIGIT mRNA expression were grouped into TMIT I, compared with other TMIT groups (Fig. 4k). Bladder cancer with TIGIT mRNA expression higher than the median value had a remarkably higher proportion of TMIT I (66.7% vs. 6.9%; P < 0.001; Fig. S4), compared with those with the fewer expression of TIGIT mRNA. More importantly, the expression of TIGIT mRNA was remarkably higher in TMIT I tumors than other TMIT groups (P < 0.001; Fig. 4l).

Finally, logistic regression analyses were performed to determine the potential statistically significant relationship between genomic characteristics and the prediction of TMIT I. The results revealed that high cytolytic activity, PD-L1 amplification, high IFN-γ mRNA expression and high TIGIT mRNA expression were independently correlated with TMIT I (Table 5).

**TMIT I was significantly associated with anti-tumor immune–related signaling pathway**

To explore the potential molecular mechanisms of TMIT I accounting for their effects on anti-PD-1/PD-L1 therapy in bladder cancer, GSEA was performed to associate the signature of genes with the TMIT I bladder tumors. The GSEA results demonstrated remarkable enrichment of signatures relating to anti-tumor immune responses, for example, the cytokine-mediated signaling pathway, IFN-γ mediated signaling pathway, activation of the immune response and MHC I-dependent antigen processing and presentation (Fig. 5).

**Discussion**

Based on the notion that effective anti-tumor immunity requires an appropriate tumor immune microenvironment, several studies have recently suggested the classification of tumors into four different types on the bases of PD-L1 expression and CD8+ CTLs statuses [6, 7, 16]. However, the underlying biology process of tumor immune microenvironment has not been fully understood until recent studies, which have shown the presence of the tumor infiltrates lymphocytes (TIL) and mutational burden are correlated with T-effector signature and immunogenic features, supporting the clinical benefits from PD-1/PD-L1 immunotherapy [11, 17, 18]. Our study was designed to identify multi-omics perspective (genomics and transcriptomics) on the tumor microenvironment on the bases of CD8 T-Cell infiltration and PD-L1 in urothelial carcinoma, and to address why TMIT I could be the potential predictive biomarker for clinical response to PD-1/PD-L1 immunotherapy from multi-omics. Our results suggested that TMIT I was proved to be correlated with high somatic mutation burden, and facilitate CD8+ T-cell infiltration and activate T-effector and IFN-γ associated gene signature. However, we did not find significantly association between TIMT subtypes and OS or DFS in urothelial cancer, which was similar to previous study in colorectal carcinomas [16]. The possible reason may be that TIMT subtypes are useful biomarkers for predicting response to PD-1 blockade. However, the urothelial cancer patients recruited in our study from the TCGA cohort did not received PD-1 blockade immunotherapy. So future clinical investigations and validations in urothelial cancer patients receiving PD-1 blockade immunotherapy are still needed.

**Table 5. Logistic regression analysis for predicting tumor microenvironment immune type I according to clinical characteristics**

|                          | Univariate analysis | Multivariate analysis | P value | P value |
|--------------------------|---------------------|-----------------------|---------|---------|
| OR (95% CI)              |                     |                       |         |         |
| Age                      | 1.003 (0.984–1.022) | 0.760                 | /       | /       |
| Gender                   | 0.821 (0.522–1.291) | 0.593                 | /       | /       |
| Number of mutations      | ≥Median 1.979 (1.314–2.981) | 0.001 | 1.055 (0.548–2.031) | 0.874 |
| Cytolytic activity       | ≥Median 33.455 (17.416–64.264) | 5.73E-26 | 5.599 (2.335–13.422) | 1.13E-04 |
| PD-L1 amplification      | Yes 4.179 (2.555–6.836) | 1.23E-08 | 11.454 (4.288–30.599) | 1.00E-06 |
| IFN-γ mRNA expression    | ≥Median 35.520 (19.100–75.704) | 6.79E-26 | 10.806 (4.453–26.224) | 1.42E-07 |
| TIGIT mRNA expression    | ≥Median 27.143 (14.461–50.251) | 8.20E-26 | 6.298 (2.702–14.671) | 2.00E-05 |

OR, odds ratio; CI, confidence interval.
The presence of neoantigen and somatic mutation burden have been proved to be associated with immunogenic features of carcinoma [9, 13]. The number of somatic mutation was considered to correlate with the chance of response to PD-1/PD-L1 blocking therapies. Here, in the TCGA cohort of bladder cancer, our linear regression analysis also found a statistically significant correlation between somatic mutation burden and the number of neoantigens. Strikingly, TMIT I group had a notably higher somatic mutation burden compared with the TMIT II tumors, indicating a more effective response to PD-1/PD-L1 blocking therapies in TMIT I bladder tumors.
Previous studies have prominently highlighted the important role of PD-L1 in immune tolerance. Urothelial carcinoma patients with PD-L1-positive cancers have a higher rate of clinical response compared to PD-L1-negative tumors [19]. In this study, the vast majority of tumors with PD-L1 amplification were classified as TMIT I. More importantly, the proportion of TMIT I in the tumors with PD-L1 amplification was remarkably higher than that in the control group without PD-L1 amplification, which indicated the potential synergistic effects of TMIT I tumors on activating PD-L1 expression. Maybe the PD-L1 amplification directly contributes to the high PD-L1 expression, which needs further validation.

The PD-L1 expression has been proved to be controlled by inflammatory signaling, produced by soluble factors from immune cells. Of those factors, IFN-γ has recently been described as the most prominent soluble inducer of PD-L1 [20]. In this study, we found the significant association between high IFN-γ mRNA expression and TMIT I. The IFN-γ mRNA expression were significantly higher in TMIT I tumors (high PD-L1 mRNA expression) than other TMIT groups, indicating potential synergistic effects on activating PD-L1 expression by IFN-γ signaling.

As the limelight novel immune checkpoint receptor target, TIGIT has been proved to be involved in mediating T cell exhaustion in tumors [21]. Blockade of TIGIT could prevent natural killer (NK) cell exhaustion and promote NK cell-dependent tumor immunity [22]. Here, we represent the first report to demonstrate the perspective of TIGIT in bladder tumor. More importantly, TIGIT mRNA expression was strikingly higher in TMIT I bladder cancer than other TMIT groups. This result is of importance because it revealed that the classification of tumors into four TMITs might be an important method for cancer immunotherapy based on other immune checkpoint inhibitors except for PD-1/PD-L1, such as TIGIT, which was associated with the dysfunction of NK cells.

Recently TP53 mutation has been considered to show significant clinical benefit to PD-1 inhibitors by altering a group of genes involved in cell-cycle regulating, DNA replication and damage repair [11]. In this study, we were surprised to found that RB1 mutation was significantly associated with TMIT I bladder cancer, and be significantly co-occurring with TP53 mutation, which could significantly activate T-effector and IFN-γ signature. However, the FGFR3 mutation was negatively correlated with TMIT I subgroup. FGFR3 and TP53 were mutually exclusive in TMIT II bladder tumors. More importantly, different amino acid changes by FGFR3/RB1 mutations were also found between TMIT I and TMIT II bladder cancer (Fig.2b, Fig.2d), such as amino acid changes in “Immunoglobulin I-set domain (260-356)” and “Protein tyrosine kinase (472-748)”, which were consistent with our GSEA observations that TMIT I was significantly associated with anti-tumor immune-related signaling pathway, including the cytokine-mediated signaling pathway, IFN-γ mediated signaling pathway and MHC I-dependent antigen processing and presentation, thereby affecting anti-tumor immune response in bladder tumors.

We detected 9 genes as significantly cancer-associated genes in TMIT I (high PD-L1/high CD8A) bladder cancer, of which, RAD51C has been reported to play an important role in DNA damage responses [23]. Remarkably, RAD51C is necessary for phosphorylation of CHK2 by ATM, which is important for checkpoint activation and acting as a favorable agent for PD-1/PD-L1 blockade therapy [24]. We also discovered different clusters of mutations in kataegis between TMIT I and TMIT II bladder tumors. Previous studies have highlighted that kataegis was characterized by clusters of C>T or C>G mutations, which were substantially enriched at TpCpN trinucleotides and on the same DNA strand [15]. It was a pity that future studies were still needed to uncover potential mechanisms that account for the different kataegis between TMIT I and TMIT II bladder tumors.

Our study has some limitations. For example, the cutoff values of CD8A and PD-L1 have not been assessed accurately. Furthermore, there were no clinical data of patients in the TCGA cohort received PD-1 blockade immunotherapy to demonstrate the real value of TMITs for response to PD-1 blockade immunotherapy in urothelial cancer directly. However, our study has some novel approaches compared to previous studies [6, 16]. Firstly, this study is ponderable since we firstly evaluated TMITs of bladder tumors based on multi-omics data. Secondly, we preliminarily investigated the underlining biology process of TMITs and found that BR1 mutation in TMIT I subset and FGFR3 mutation in TMIT II subset might play important roles in predicting clinical response to PD-L1/PD-1 immunotherapy. And thirdly, we represented the first report to demonstrate the perspective of TIGIT in bladder tumor, which could prevent NK cell exhaustion and promote NK cell-dependent tumor immunity.

Conclusion

In summary, our study provided a multi-omics perspective on the tumor microenvironment on the
bases of CD8 T-Cell infiltration and PD-L1 in urothelial carcinoma. The classification of bladder cancer into four TMITs on the bases of the PD-L1 expression and the CD8+ CTLs statuses is an appropriate approach for bladder tumor immunotherapy. There was not significantly association between TMIT subtypes and OS or DFS in urothelial cancer. TMIT I (high PD-L1/high CD8A) is significantly correlated with more somatic mutation burden, and facilitates CD8+ T-cell infiltration and activates T-effector and IFN-γ associated gene signature. Alteration landscape for somatic variants was different between TMIT I and TMIT II (low PD-L1/low CD8A).

Abbreviations

TCGA: The Cancer Genome Atlas; PD-1: programmed cell death 1; PD-L1: PD-1 ligand; CTL: cytotoxic T cells; TMIT: tumor microenvironment immune type; MAF: mutation annotation format; CAN: copy number alteration; GSEA: Gene Set Enrichment Analysis; FDR: false discovery rate; NES: normalized enrichment score; Ti/Tv: transition/transversion; IFN-γ: interferon gamma; TIL: tumor infiltrating lymphocytes.

Supplementary Material

Supplementary figures. http://www.jcancer.org/v10p0697s1.pdf

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Competing Interests

The authors have declared that no competing interest exists.

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