Metabolic-associated signature and hub genes associated with immune microenvironment and prognosis in bladder cancer

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Abstract
The relationship between metabolism and immune microenvironment remains to be studied in bladder cancer (BCa). We aimed to construct a metabolic-associated signature for prognostic prediction and investigate its relationship with the immune microenvironment in BCa. The RNA expression of metabolism associated genes was obtained from a combined data set including The Cancer Genome Atlas, GSE48075, and GSE13507 to divide BCa patients into different clusters. A metabolic-associated signature was constructed using the differentially expressed genes between clusters in the combined data set and validated in the IMvigor210 trial and our center. The composition of tumor-infiltrating immune cells (TIICs) was evaluated using the single-sample Gene Set Variation Analysis. BCa patients in Cluster A or high-risk level were associated with advanced clinicopathological features and poor survival outcomes. The percentage of high-risk patients was significantly lower in patients responding to anti-PD-L1 treatment. Compared with low-risk patients, the IC50 values of cisplatin and gemcitabine were significantly lower in high-risk patients. Thiosulfate transferase (TST) and S100A16 were significantly associated with clinicopathological features and prognosis. Downregulation of TST promoted BCa cell invasion, migration, and epithelial-to-mesenchymal transition, which are inhibited by downregulation of S100A16. CD8 + T cells, neutrophils, and dendritic cells had higher infiltration in the TST low-level and the S100A16 high-level. Furthermore, loss of function TST and S100A16 significantly affected the expression of PD-L1 and CD47. The metabolic-associated signature can stratify BCa patients into distinct risk levels with different immunotherapeutic susceptibility and survival outcomes. Metabolism disorder promoted the dysregulation of immune microenvironment, thus contributing to immunosuppression.

KEYWORDS
bladder cancer, immune microenvironment, metabolism, prognosis, risk signature

Abbreviations: BCa, bladder cancer; BCG, Bacillus CalmetteGuérin; CCLE, cancer cell line encyclopedia; CR, complete response; DEGs, differentially expressed genes; DSS, disease-specific survival; EMT, epithelial-to-mesenchymal transition; GSEA, gene set enrichment analysis; HR, hazard ratio; IHC, Immunohistochemistry; MAGs, metabolism associated genes; MMP9, Matrix metalloproteinase 9; NMIBC, nonmuscle-invasive bladder cancer; PD, progressive disease; PR, partial response; RT-qPCR, real-time quantitative PCR; S100A16, S100 calcium-binding protein A16; SD, stable disease; ssGSEA, single-sample gene set enrichment analysis; TAMS, tumor-associated macrophages; TCGA, The Cancer Genome Atlas; TIICs, tumor-infiltrating immune cells; Tregs, regulatory T cells; TST, thiosulfate transferase; TURBT, transurethral resection of bladder tumor.
1  |  INTRODUCTION

Bladder cancer (BCa) ranks the 10th most common malignancy in the world, with approximately 17,980 deaths and 81,400 new cases in the USA in 2020.1 BCa can be classified into muscle-invasive bladder cancer (MIBC, 25%) and nonmuscle-invasive bladder cancer (NMIBC, 75%).2 NMIBC patients typically receive the transurethral resection of bladder tumor followed by intravesical therapy with Bacillus CalmetteGuérin for patients with carcinoma in situ or high-grade T1.3 As for MIBC patients, radical cystectomy or chemotherapy is typically recommended.4,5 To date, the treatment options for BCa patients are still limited, and the rates of recurrence, progression and metastasis are still disappointingly high.6,7 In addition, the treatment strategy and prognosis prediction mainly rely on tumor staging system, which is not sufficient to accurately predict the prognosis and develop the personalized treatment in BCa.8 Therefore, exploration of effective biomarkers for prognosis prediction and clinical management is still challenging.

Dysregulations of the metabolism and immune microenvironment within tumor are key events in various tumors.9,10 Previous studies have reported that the dysregulation of some metabolic pathways, including activation of fatty acid metabolism, glutamine metabolism and glycolysis, are highly related to the tumorigenesis and tumor progression in BCa.11,12 In addition, studies have predicted the prognosis of BCa patients and treatment response by metabolic classification, which indicates that metabolism plays an important role in BCa.13 Thus, exploration of metabolism associated genes (MAGs) may be useful in discovering the key metabolic-associated mechanisms controlling carcinogenesis, which may facilitate the development of new treatment options. Furthermore, some studies have found important effects of metabolism on immune cells in tumors, which lead to the tumor microenvironment being generally acidic, hypoxia and/or depletion of key nutrients needed by immune cells.14 It has been reported that tumor metabolites affect the function of tumor-associated macrophages (TAMs), and TAMs further activate glycolysis, fatty acid synthesis, and alter nitrogen cycle metabolism by reprogramming their metabolism to promote tumor progression and metastasis.15 Moreover, macrophage subtypes associated with immunosuppression, tumor angiogenesis, and progression affect the progression of BCa.16 However, no systematic assessment of the association between metabolism phenotype and immune microenvironment (including tumor-infiltrating immune cells [TIICs] and the expression of critical immune checkpoint genes) has been performed in BCa.

In this study, we used the transcriptomic data to establish and validate a metabolic-associated signature for prognostic prediction. In addition, we further investigated the relationship between the metabolic-associated signature/hub genes and the immune microenvironment.

2  |  MATERIALS AND METHODS

2.1  |  Ethics

This study was reviewed and approved by the Ethics Committee of Shanghai Tenth People’s Hospital, and all patients signed written informed consent.

2.2  |  Data collection

The list of MAGs was extracted from the metabolic pathways Gene set enrichment analysis (GSEA, c2.cp.kegg.v7.1.symbols.gmt). We used the ComBat, a batch effect correcting tool in the sva R package v3.38, to adjust batch effects and combine these data sets.

2.3  |  Unsupervised clustering

We used the “ConsensusClusterPlus” R package v.1.50.0 (Pearson correlation, 50 iterations and 80% resample rate) to divide BCa patients into different clusters based on the expression of MAGs. Gene Set Variation Analysis (GSVA) was performed to quantitatively investigate different biological pathways between clusters using “GSVA” R package v.1.38.2. single-sample gene set enrichment analysis (ssGSEA) was used to evaluate the proportions of 28 TIICs in each BCa sample.

2.4  |  Construction of the metabolic-associated signature

The differentially expressed genes (DEGs, |Log2FC| > 1.5 and adjusted p < 0.05) between clusters were identified using "limma" R package v3.46. The random forest along with Boruta feature selection was performed using the “Boruta” R package v7.0 to obtain the most important DEGs involved in discriminating clusters, followed by the univariate Cox regression. In this way, the selected genes can be divided into two groups based on the hazard ratio (HR) of the univariate Cox regression: protective gene (HR < 1) and risk gene (HR > 1). The metabolic-associated signature was developed using the following formula:

metabolic score= (scale ∑X − ∑Y).

Where X is the expression of each selected gene with HR > 1, and Y is the expression of each selected gene with HR < 1. This formula was used to calculate the risk score of each patient. Then, patients were divided into low- and high-risk levels according to the optimal cutoff value of risk score calculated by the “survminer” R package v0.4.8.
2.5 | Chemotherapeutic response prediction

The chemotherapy response of each sample was predicted based on the Genomics of Drug Sensitivity in Cancer which is the largest public pharmacogenomics database. Two commonly used chemicals Cisplatin and Gemcitabine were selected to evaluated the IC50 of each sample using the “pPRoPhetic” R package.

2.6 | Gene mutation in cancer cell line encyclopedia (CCLE)

The gene expression data and mutation data of 592 carcinoma cell lines were down from CCLE [https://portals.broadinstitute.org/ccle]. The risk scores of each carcinoma cell line were calculated using the formula mentioned above. Then, carcinoma cell lines were divided into low- and high-risk levels based on the median value of risk scores. The mutation counts were compared between low- and high-risk levels.

2.7 | Hub genes screening

We used three algorithms, including support vector machine-based recursive feature elimination (SVM-RFE, “e1071” R package v1.7) algorithm, MeanDecreaseGini-based random forest-feature selection (RFS-FS, “randomForest” R package v4.6) and MeanDecreaseAccuracy-based RFS-FS, to rank genes with respect to the risk level. The top 10 genes of each algorithm were put into a Venn online software (http://bioinformatics.psb.ugent.be/webtools/Venn/) to identify the overlapping hub genes. The GSEA (http://www.broadinstitute.org/GSEA) was used to investigate the enriched gene signatures.

2.8 | Bioinformatics analysis

The Assistant for Clinical Bioinformatics is an online tool (www.aclbi.com) that contains gene expression data and survival information of various cancers in the TCGA database, and all data are analyzed by R v4.0.3. This data set can provide survival analysis, correlation analysis, prognostic nomogram and tumor tissue immune cell infiltration.

Kaplan–Meier Plotter is an online tool (www.kmplot.com) that contains gene expression data and survival information of various cancers for analyzing the prognostic value of thiosulfate transferase (TST) and S100A16 in BCa. According to the median mRNA expression of BCa samples, they were divided into high expression level and low expression level, and then analyzed by Kaplan–Meier survival curve.

TIMER is an online tool (http://timer.cistrome.org) that contains a comprehensive analysis of immune infiltration in different types of cancer, and is used to analyze the association between TST and S100A16 and immune infiltration in BCa. The associations between genetic markers and various immune cells are also recorded.

GEPIA is an online tool (http://gepia2.cancer-pku.cn) that can provide tumor/normal differential expression analysis, survival analysis, correlation analysis. BCa patients were used to analyze the association between genes based on their mRNA expression.

2.9 | Immunohistochemistry (IHC)

The Human Protein Atlas (http://www.proteinatlas.org) online database can obtain IHC slides and information to check the expression levels of TST and S100A16 in normal bladder tissues and BCa specimens. The score of the positive staining ratio of tumor cells is defined as: 0 (none); 1 (<25%); 2 (25%~75%); 3 (>75%). The staining intensity score is determined as: 0 (negative); 1 (weak); 2 (moderate) and 3 (strong). The immune response score is evaluated by multiplying the staining intensity and the positive cell ratio score.

2.10 | Cell lines and culture

Human HEK293T, human BCa cell line T24 were obtained from the American Type Culture Collection (ATCC). Human HEK293T is maintained in DMEM medium (Gibco); T24 cells are cultured in RPMI-1640 (Gibco). All cell lines were cultured in a humidified incubator containing 5% CO₂ at 37°C. All cell culture media contain 10% fetal bovine serum (FBS; Gibco) and 1% penicillin/streptomycin (Hyclone; GE Healthcare Life Sciences).

2.11 | Generation of TST and S100A16 knockout cell lines

TST and S100A16 gene knockout T24 cell lines were produced using lentivirus CRISPR technology. In short, the oligonucleotides encoding gRNA (TST sgRNA1: 5’-TCCGGAACTGGCTGAAGGA-3’; TST sgRNA2: 5’-CATCAGGACTGGCAAGCTG-3’; S100A16 sgRNA1: 5’-AGGCCCTTACCCAGCACATG-3’; S100A16 sgRNA3: 5’-CAGGACACGGGAACCGGA-3’) was constructed into the lentiviral CRISPR-puro vector. Then, the plasmid was cotransfected into HEK293T cells with packaging vectors including pSPAX2 and pMD2G. After 48 h, the culture supernatant was harvested to infect the T24 cell line. Puromycin (1 μg/ml) was used to select lentivirus-transduced cells for 3 days and verified by Western blot.

2.12 | Real-time quantitative PCR (RT-qPCR)

According to the manufacturer’s instructions, total RNA was extracted from the cells using TRIzol reagent (Invitrogen), and then reverse transcription was performed using HiScript®Q Select RT SuperMix (Vazyme Biotechnology Company) to synthesize...
2.13 | Western blot

The cells were collected and lysed in lysis buffer (150 mM NaCl, 0.5% NP-40, 1 mM EDTA, 10% glycerophosphate, 50 mM Tris-Cl, pH 7.4 and protease inhibitor cocktail). After 30 min, the cell lysate was separated by centrifugation at 12,000 rpm 4°C for 15 min. The protein concentration was quantified by BCA protein electrophoresis (SDS-PAGE), and analyzed by immunoblotting. The western blot was detected using an electrochemiluminescence imaging system (Tanon). Anti-TST (ab166625), CD47 (ab218810), PD-L1 (ab136845), E-cadherin (ab1416), N-cadherin (ab12221), Vimentin (ab92547), Slug (ab85936) antibodies were obtained from Abcam; anti-Tubullin (2148S) antibody was obtained from Cell Signaling Technology; anti-β‐S100A16 (114561-AP) antibody was obtained from Proteintech.

2.14 | Flow cytometry

Cells were washed in PBS and stained for cell surface CD47 or PD-L1 on ice for 30 min in PBS plus 2% FACS. After washing in PBS samples were analyzed on a BD LSR Fortessa and analyzed in FlowJo. APC antihuman CD274 Antibody Obtained from Biolegend, Clone 29E.2A3, Cat# 329708; PE antihuman CD47 Antibody Obtained from Biolegend, Clone: CC2C6, Cat# 323108.

2.15 | Cell migration assay

According to the manufacturer's instructions, cell migration was analyzed using the Transwell system (Corning). Resuspend the T24 cell line in serum-free medium, and then add 200 μl of cell suspension (containing $5 \times 10^4$ cells) to the upper Transwell chamber. The lower chamber contains 600 μl of medium with 10% FBS. After 16 h of incubation at 37°C, the cells that migrated on the lower surface of the chamber were fixed with 4% paraformaldehyde and stained with crystal violet.

2.16 | Matrigel droplet invasion assay

$5 \times 10^4$ T24 cells were suspended in 10 μl Matrigel, seeded in a 24-well plate in the form of droplets, and the droplets were imaged using a microscope (Olympus Corporation) on Day 0 and Day 4, as described previously. The added medium was changed once a day. The migration of cells extending to the outside of the droplet was measured by the ImageJ software.

2.17 | Statistical analysis

Statistical analysis was conducted with SPSS 23.0 (IBM Corp.) and R statistical software v3.6.1 (https://www.r-project.org/). The factors between clusters or risk levels were compared using the $\chi^2$ test, the t-test or the Mann–Whitney U-test, as appropriate. Survival analyses were performed using the Kaplan–Meier method and Cox proportional hazard models. All tests were 2-tailed, and $p < 0.05$ were considered statistically significant.

3 | RESULTS

3.1 | Consensus clustering identified two clusters of BCa patients

The expression of 1257 MAGs were achieved from TCGA, GSE48075 and GSE13507. These three data sets (712 patients) were combined for clustering. We divided the combined data set into several clusters based on the consensus of expression of 1257 MAGs (Figure 1). When the “k” (clustering index) increased from 2 to 9, $k = 2$ was the optimal value to obtain the minimal interference (Figure 2A) and largest differences (Figure 2B) between clusters. In this way, the combined data set was classified into two robust clusters, namely Cluster A and Cluster B. Kaplan–Meier survival analysis revealed that Cluster A had a significantly poor OS than Cluster B (Figure 2C). Cluster A was significantly associated with advanced clinicopathological features, including tumor stage, tumor grade, histological subtype, and age (Figure 2D). GSEA demonstrated that infection-related biological processes and metabolic-related biological processes were mainly enriched in Cluster A and Cluster B, respectively (Figure 2E). ssGSEA showed that the proportions of 27 TIICs were different between Cluster A and B (Figure 2F).

3.2 | Development of the metabolic-associated signature

There were 1721 DEGs between Cluster A and B. After the feature filtering by the random forest and the univariate cox regression, 159 genes (HR > 1: 56 genes, HR < 1: 103 genes; Supporting Information: Table 2) were selected for the development of the metabolic-associated signature using the formula mentioned above. In the combined data set, BCa patients were divided into low- and high-risk levels based on the optimal cutoff value (~0.03655953). Patients in the high-risk level had significantly poor survival outcomes than patients in the low-risk level (Figure 3A). In addition, the IC50 values of Cisplatin and Gemcitabine were significantly higher in patients with low-risk scores (Figure 3B,C).
In TCGA, the proportions of 26 TIICs were different between low- and high-risk levels (Figure 3D), and patients with malignant clinicopathological features, including nonpapillary subtype, high tumor grade, high tumor stage, older in age (≥60 years), female and smoking history, had significantly higher risk scores (Figure 3E). In addition, patients with high-risk scores exhibited poor OS compared with those with low-risk scores (Figure 3F).

3.3 | Genetic alteration in CCLE

592 carcinoma cell lines were divided into low- and high-risk levels based on the median cutoff value (~0.06701084). The top 20 mutation genes in low- and high-risk levels were presented in Supporting Information: Figure 1, which revealed the difference in genetic alteration between low- and high-risk levels. In addition, carcinoma cell lines with low-risk scores had significantly higher mutation than those with high-risk scores (Figure 3G).

3.4 | Validation of the metabolic-associated signature

In IMvigor210 trial, patients treated with PD-L1 inhibitor were divided into low- and high-risk levels based on the optimal cutoff value (0.6055458). Kaplan–Meier survival analysis revealed that high-risk patients had significantly poor OS than low-risk patients (Figure 4A). In addition, the percentage of high-risk patients was significantly lower in patients responding to anti-PD-L1 treatment (complete response [CR]/partial response [PR]) than in patients nonresponding to anti-PD-L1 treatment (progressive disease [PD]/stable disease [SD]) level (Figure 4B), which indicating the potential
clinical value of the metabolic-associated signature to identify the immunotherapeutic susceptibility for patients with BCA.

In our center a total of 135 BCa samples (108 male, 27 female; mean age = 70.4 years ± 11.7 [standard deviation], range 31–95 years; 99 NMIBC, 36 MIBC; 30 low-grade, 105 high-grade) were collected for RNA-sequence. High-grade patients and MIBC patients had significantly higher risk scores than NMIBC patients and low-grade patients, respectively ($p < 0.001$ and $p = 0.003$, Figure 4C,D).
FIGURE 3  Performance of the metabolic-associated signature. (A) Kaplan–Meier overall survival curves of the two risk levels. (B) Cisplatin IC50 of the two risk levels. (C) Gemcitabine IC50 of the two risk levels. (D) Tumor-infiltrating immune cells of the two risk levels. (E) Clinicopathological features of the two risk levels in The Cancer Genome Atlas (TCGA). (F) Kaplan–Meier overall survival curves of the two risk levels in TCGA. (G) Mutation count of the two risk levels in Cancer Cell Line Encyclopedia. *p < 0.05, **p < 0.01, ***p < 0.001. [Color figure can be viewed at wileyonlinelibrary.com]
3.5 | The prognostic value of hub genes

Through hub genes screening, three genes, including TST, S100 calcium binding protein A16 (S100A16) and Matrix metalloproteinase 9 (MMP9) were obtained (Figure 5A). The functions of MMP9 have been fully investigated in various tumors, so the experimental validation of TST and S100A16 was performed in this study. Validation in the Human Protein Atlas shows that the protein levels of TST and S100A16 in BCa are different from those in normal bladder tissues, especially S100A16 (Figure 5B,C). In addition, RNA-sequence data in our center showed that the expressions of TST and S100A16 were significantly correlated with T stage (Figure 5D) and histopathological grade (Figure 4E). The Assistant for Clinical Bioinformatics and Kaplan–Meier plotter database analysis showed that the low expression level of TST and the high expression level of S100A16 predicted poor disease-specific survival (DSS) and OS of BCa patients (all \( p < 0.05 \), Figure 5F,G).

Univariate Cox proportional hazards regression analysis showed that age, pTNM stage, low expression of TST and high expression of S100A16 were associated with poor OS of BCa patients in TCGA (Figure 6A). Then, multivariate Cox regression analysis showed that age, pTNM stage, TST and S100A16 were independent prognostic factors of BCa patients (Figure 6B). Then, we combined age, TST and S100A16 to generate a prognosis nomogram. The results showed that the nomogram could usefully predict the 1-, 3-, and 5-year survival outcomes of BCa patients (Figure 6C,D). Taken together, these findings indicate that TST and S100A16 are significantly related to the clinicopathological features and prognosis of BCa patients, which indicates the importance of further research.

3.6 | Downregulation of TST promoted the invasion, migration and epithelial-to-mesenchymal transition (EMT) of BCa cells, while downregulation of S100A16 inhibited those processes

We synthesized two specific sgRNAs for TST and S100A16 respectively, and studied whether TST and S100A16 affected the migration and invasion of BCa using matrigel drop infiltration assay and transwell migration assay. First, we confirmed by western blot
FIGURE 5  Validation of hub genes in metabolic-associated signature. (A) thiosulfate transferase (TST), S100A16 and MMP9 were obtained by hub genes screening. (B) IHC of TST and S100A16 in human protein atlas. (C) IHC-based H score of TST and S100A16. (D, E) The expression of TST and S100A16 in different T stages (D) and Grade (E). (F, G) Overall survival and disease-specific survival curves in TST (F) and S100A16 (G) high and low expression bladder cancer cases from the Assistant for Clinical Bioinformatics (www.aclbi.com) and Kaplan–Meier plotter database (http://kmplot.com/analysis/). Error bar ± SD; *p < 0.05. [Color figure can be viewed at wileyonlinelibrary.com]
that the expression of TST and S100A16 in T24 cells can be effectively knocked out by sgRNA (Figure 7A). Then matrigel droplet measurement was performed, and the radial distance of cells moving out from the edge of matrix glue droplet was measured on the 4th day after spreading the cells. Compared with the control level, the downregulation of TST enhanced the invasion ability of BCa cells, while downregulation of S100A16 inhibited these processes (Figure 7B–E). Similarly, we measured the migration of T24 cell by transwell migration assay, which was consistent with the above results. The results showed that downregulation of TST enhanced the invasion and migration of BCa cells, while downregulation of S100A16 inhibited those processes (Figure 7F–I).

Next, to explore the detailed mechanism of TST and S100A16, we used GSEA to analyze the TST and S100A16 related signal pathways of BCa in TCGA database. GSEA analysis showed that TST-low level and S100A16-high level were significantly positive in FOCAL ADHESION, GAP JUNCTION (Figure 7J,K) and ADHERENS JUNCTION, which means that TST and S100A16 may play a key role in the progression of tumor EMT. Based on this, the protein expression level of EMT markers were detected. These results showed that E-cadherin decreased, while mesenchymal markers including N-cadherin, vimentin and slug increased in T24 cell line in which knocked out TST (Figure 6L). Conversely, E-cadherin was upregulated and N-cadherin, vimentin and slug were downregulated in T24 cell line which knocked out S100A16 (Figure 6M). These results indicate that TST and S100A16 play a vital role in invasion, migration and EMT of BCa cells.

### 3.7 Correlation between TST and S100A16 and tumor immune microenvironment of BCa

GSEA analysis showed that the gene sets of BLADDER CANCER, ANTIGEN PROCESSING AND PRESENTATION, NATURAL KILLER CELL MEDIATED CYTOTOXICITY, CYTOSOLIC DNA SENSING PATHWAY were significantly enriched in patients with lower levels...
of TST expression and higher levels of S100A16 (Figure 8A,B; Supporting Information: Figure 2A,B). The results of the Assistant for Clinical Bioinformatics showed that CD8+ T cells, neutrophils and dendritic cells had higher immune infiltration in the TST-low level and S100A16-high level (Supporting Information: Figure 2C,D).

And then, the correlation between TST, S100A16 and immune cell infiltration and the influence of different immune cells on the prognosis of BCa were analyzed by using TIMER database. According to the results of TIMER, the expression of TST was negatively correlated with the infiltration of CD8⁺ T cells, CD4⁺ T cells, neutrophils and dendritic cells, while the expression of S100A16 was positively correlated with these immune cells (p < 0.05, Supporting Information: Figure 2E,F). TIMER for “survival” module showed that the high expression of CD8⁺ T cells and S100A16 had worse
FIGURE 8  Correlation between TST and S100A16 and tumor immune microenvironment of BCa. (A, B) GSEA analysis of TST (A) and S100A16 (B). (C, D) The total protein and mRNA expression levels of PD-L1 and CD47 were detected in TST (C) and S100A16 (D) knocked out T24 cells, respectively. (E–H) The membrane protein expression level of PD-L1 and CD47 were detected in TST (E, F) and S100A16 (G, H) knocked out T24 cells, respectively. (I–K) TCGA database showed that TST was negatively correlated with PDL1 (I) and CD47 (J), and S100A16 was positively correlated with CD47 (K). BCa, bladder cancer; GSEA, Gene set enrichment analysis; mRNA, Messenger RNA; TCGA, The Cancer Genome Atlas; TST, thiosulfate transferase. [Color figure can be viewed at wileyonlinelibrary.com]
prognosis, while patients with high expression of TST had better prognosis (all p < 0.05, Supporting Information: Figure 2G).

Furthermore, we analyzed the correlation between TST and S100A16 in BCa and immune-related checkpoints in TCGA. Our results showed that TST and S100A16 were significantly correlated with most immune-related checkpoints, especially PD-L1 and CD47 (p < 0.05, Supporting Information: Figure 2H,I). To test whether TST and S100A16 really participate in the regulation of PDL1 and CD47, we transfected T24 cell with TST and S100A16 specific Cas9/sgRNA, and S100A16 really participate in the regulation of PDL1 and CD47, TST knockout T24 cell (Figure 8C,E,F), and TST was significantly increased in mRNA, total protein and membrane protein levels. Our data showed that PD-L1 and CD47 were increased in mRNA, total protein and membrane protein levels in TST knockout T24 cell (Figure 8C,F), and TST was significantly negatively correlated with PD-L1 and CD47 (Figure 8I,J). In addition, CD47 increased in mRNA, total protein and membrane protein levels in S100A16 knockout T24 cells (Figure 8D,G), and S100A16 was positively correlated with CD47 (Figure 7K), while PD-L1 did not change (Figure 8D,H). To sum up, these findings indicate that TST and S100A16 play a key role in the regulation of tumor immune microenvironment of BCa.

4 | DISCUSSION

Metabolic dysregulation is emerging as a novel hallmark in tumor and plays important role in tumorigenesis and tumor progression. Previous studies had reported that the fatty acid metabolism was highly activated in BCa tissues compared with paracancerous tissues, and the alteration in glucose and pyruvate metabolism could promote the progression of BCa cells. Considering the vital role of the metabolic disorder in BCa, it is important to identify metabolic-related prognostic biomarkers for the prognosis and treatment of BCa. In this study, we identified two clusters based on the RNA expression of 1257 MAGs. The clinicopathological features and prognoses were significantly different between the two clusters, revealing that the RNA expression of 1257 MAGs remarkably related to features of BCa. GSVA demonstrated that clusters A and B were primarily enriched for infection- and metabolism-related biological processes, respectively, which may partly explain the differences in clinical pathology and prognosis between the two clusters. However, according to their metabolic classification, tang et al. found that the biological processes of the three clusters were all significantly related to the metabolic pathways. The reason for this difference may be due to inconsistent algorithms for metabolic classification.

Then, we used the hub DEGs between two clusters to construct a metabolic-associated signature that had good performance in prognostic stratification in the combined data set and IMvigor210 trial, in which BCa patients with high-risk scores showed significantly poorer survival outcomes than patients with low-risk scores. Moreover, patients with high-risk scores were more prone to have advanced clinicopathological features and have SD or PD when receiving PD-L1 inhibitors, which revealed that patients with low-risk scores are more to respond to immunotherapy. In addition, the IC50 values of Cisplatin and Gemcitabine were significantly lower in patients with high-risk scores, which revealed that patients with high-risk scores are more to respond to chemotherapy. Based on the above results, it may be helpful to guide the individualized treatment of BCa. Tang et al. also predicted the treatment response of patients with BCa by metabolic classification. Their study found that patients with M1 subtype showed the best predictive response to immunotherapy and were the least sensitive to cisplatin. By contrast, patients with M2 subtypes had the worst predictive response to immunotherapy, but were more sensitive to chemotherapeutic agents such as cisplatin and doxorubicin. Similar conclusions were reached, although a different algorithm was used as in our study.

High needs of cancer cells for nutrients contribute to the metabolic reprogramming of both cancer cells and TIICs, which may shape the density/composition of TIICs, or directly modulate the functionality of TIICs to facilitate immunosuppression and tumor progression. TAMs and regulatory T cells (Tregs) represent the main fraction of the TIICs in the tumor microenvironment to support tumor growth and metastasis through promoting immunosuppression. It has been reported that the activation of fatty acid β-oxidation in TAMs and regulatory T cells is important for their survival and immunosuppression function in tumor progression. In our study, the composition of TIICs between clusters or risk levels was distinct, and patients in cluster A or high-risk level had significantly higher proportions of TAMs and regulatory T cells, which may contribute to the poor survival outcomes of these patients. In addition, low mutation count and high proportion of TAMs and regulatory T cells in high-risk level partly explained the low proportion of CR/PR and poor survival outcomes of high-risk patients treated with PD-L1 inhibitor. Similarly, Tang et al. found a positive correlation between high immune infiltration and poor therapeutic response of immune checkpoint inhibitors in the prediction of treatment response. Thus, the metabolic disorder could shape the composition of TIICs and the metabolic-associated signature may have the potential to facilitate personalized immunotherapy in BCa.

Experimental studies of hub genes in the metabolic-associated signature were performed to explore their immunological effects. TST is an abundant mitochondrial enzyme that catalyzes the transfer of sulfur in multiple biological pathways, especially the conversion of sulfonated Glutathione to thiosulfate and glutathione in a sulfite-dependent manner. Dysfunction of TST was highly associated with the development of metabolic diseases such as obesity and diabetes. It is reported that the TST may play a role in the development of colorectal cancer and glioma. S100A16 belongs to the small acidic S100 calcium-binding protein family that is involved in chromosomal instability and rearrangements, contributing to malignant transformation of cells. Overexpression of S100A16 in diverse cancer cells reveals its close relationship with the tumorigenesis and tumor progression. However, the functions of these two genes in BCa cells have not been explored. In this study, we explored the biological functions and immunological effects of TST and S100A16 in BCa cells. The results demonstrated that TST...
knockdown promoted the invasion, migration and EMT of the BCa cell while S100A16 knockdown inhibited these processes. Studies have shown that TST can detoxify H$_2$S,$^{32}$ and H$_2$S is crucial for the occurrence and progression of BCa, especially in inducing cell proliferation and promoting invasion.$^{40-42}$ Similarly, some studies have found that S100A16 promotes drug resistance in NMIBC,$^{43}$ and S100A16 also promotes the metastasis and progression of pancreatic cancer by inducing EMT.$^{44}$ In addition, the results of western blot and flow cytometry showed that TST knockdown promoted and S100A16 knockdown inhibited the protein expressions of critical immune checkpoint genes that have functions in immunosuppression. Thus, dysexpression of TST or S100A16 may contribute to the tumor progression, immunosuppression and dysregulation of immune microenvironment in BCa.

Despite the above important findings, our study has certain limitations. First, this is a retrospective study, which may lead to deviations due to incomplete information. Second, the sample size of the model needs to be further expanded before clinical application. Finally, additional functional studies and mechanism analysis are still needed to explore the potential molecular mechanisms of the immune effects of TST and S100A16, which are of great significance for the development of targeted therapy.

5 | CONCLUSION

The metabolic-associated signature had the potential to help outcome prediction and personalized immunotherapy in BCa. Metabolism disorder contributed to the dysregulation of immune microenvironment, thus leading to immunosuppression. The hub genes in the metabolic-associated signature could modulate the tumor immune microenvironment.

AUTHOR CONTRIBUTIONS

Zongtai Zheng and Yadong Guo collected and analyzed research data. Shenghua Liu was responsible for the interpretation of the results. Yadong Guo and Zongtai Zheng drafted the manuscript. Shiyu Mao, Fuhan Yang, Ruiliang Wang, Hong Wang, Cheng Li, Ji Liu, Qinwan Wang, Xudong Yao, and Shenghua Liu performed the study design and revised the manuscript. All authors have read and agreed to the published version of the manuscript.

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CONFLICT OF INTEREST

The authors declare no conflicts of interest. A preprint was submitted in Research Square at the link: https://www.researchsquare.com/article/rs-1041335/v1

DATA AVAILABILITY STATEMENT

We collected 135 BCa samples confirmed histologically in Shanghai Tenth People’s Hospital between November 2019 and April 2021 according to the following criteria: (1) histologically confirmed BCa; (2) samples were available for RNA-sequence; (3) availability of clinical and follow-up data. The processes of total RNA extraction, paired-end libraries generation and RNA-sequence can be found in our previous publications.$^{16}$ RNA-sequence data (FPKM standardized data) for BCa patients were downloaded from The Cancer Genome Atlas (TCGA) (http://www.cancergenome.nih.gov). The microarray data sets GSE48075 and GSE13507 were obtained from the GEO database (https://www.ncbi.nlm.nih.gov/). The transcriptomic data of IMvigor210 trial, which included metastatic urothelial cancer patients treated with atezolizumab (PD-L1 inhibitor), were downloaded from the http://research-pub.gene.com/IMvigor210CoreBiologies

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SUPPORTING INFORMATION
Additional supporting information can be found online in the Supporting Information section at the end of this article.

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