Construction and validation of a novel prognostic signature for uveal melanoma based on five metabolism-related genes

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Abstract: Background: Uveal melanoma (UM) is the most aggressive intraocular tumor worldwide. Accurate prognostic models are urgently needed. The present research aimed to construct and validate a prognostic signature is associated with overall survival (OS) for UM patients based on metabolism-related genes (MRGs). Methods: MRGs were obtained from molecular signature database (MSigDB). The gene expression profiles and patient clinical data were downloaded from The Cancer Genome Atlas (TCGA) database. In the training datasets, MRGs were analyzed through univariate Cox regression analyses and least absolute shrinkage and selection operator (LASSO) Cox analyses to build a prognostic model. The GSE84976 was treated as the validation cohort. In addition, time-dependent receiver operating characteristic (ROC) and Kaplan-Meier survival curve analyses the reliability of the developed model. Then, gene set enrichment analysis (GSEA) was used for gene enrichment analysis. Nomogram that combined the five-gene signature was used to evaluate the predictive OS value of UM patients. Results: Five MRGs were identified and used to establish the prognostic model for UM patients. The model was successfully validated using the testing cohort. Moreover, ROC analysis demonstrated a strong predictive ability that our prognostic signature had for UM prognosis.
Multivariable Cox regression analysis revealed that the risk model was an independent predictor of prognosis. UM patients with a high-risk score showed a higher level of immune checkpoint molecules. **Conclusion:** We established a novel metabolism-related signature that could predict survival and might be therapeutic targets for the treatment of UM patients.

**Keywords:** uveal melanoma; metabolism; TCGA; GEO; prognostic model; immune cell infiltration

1. Introduction

Uveal melanoma (UM) is the most aggressive intraocular malignant tumor in adults and has a poor prognosis. The mean age-adjusted incidence is 5.1 per million in the United States [1]. UM is considered a high molecular heterogeneous and immunogenicity disease [2]. The progression of clinical treatment includes surgery, photocoagulation, and radiotherapy, which can achieve great local tumor control [3]. Nevertheless, previous studies have reported that up to 50% of UM patients will have experienced a local tumor recurrence by 10 years, and the survival rate remains poor [4,5]. As UM has the tendency to early metastasis, nearly 50% of UM patients develop tumor metastasis via hematogenous spread primarily to the liver [1]. The metastatic UM patients are usually fatal within 1 year of the onset of symptoms [6]. During the past decades, immune checkpoint blockade (ICB) has been proven efficacious in cutaneous melanoma. Unlike cutaneous melanoma metastases, UM metastases generally respond poorly to ICB, with a response rate of only 0–5% [7]. Despite being treated for their primary intraocular tumor, these patients are at lifetime risk of developing metastasis. Therefore, it is crucial to reveal a new biomarker to assess UM prognosis.

Dysregulated metabolism is one of the well-recognized hallmarks of solid human tumors. It relates to the rapid proliferation and preferential survival of cancer cells [8]. During tumorigenesis and metastasis, aberrantly activated nutrient acquisition metabolism pathways facilitate the reprogramming of cancer cell metabolism to meet the bioenergetic, biosynthetic, and redox demands of malignant cells [9]. Otto Warburg first observed the reprogrammed metabolic pathway in the cancer cell, which was later termed the “Warburg effect” or “aerobic glycolysis” to describe that cancer cells take up glucose and produce lactate regardless of oxygen availability [10]. Previous studies have reported a critical metabolic program within poor prognostic monosomy 3 UM, associated with higher metastasis and poor prognosis [11]. Monosomy 3 UM exhibited a less glycogenetic and more insulin-resistant phenotype, together with reduced glycogen levels, which were associated with the development of metastases and a reduced survival rate [12]. UM has the highest oxidative phosphorylation characteristics among 31 tumor types. A recent study also indicates that high cysteinyl leukotriene receptors, a lipid-signaling molecule, expression correlates with poor survival in UM patients [13]. In addition, some energy metabolism protein levels were significantly different in glycolysis after irradiation [14]. However, the relationship between metabolism-related genes (MRGzs) and UM prognosis has not been fully elucidated.

With the development of next-generation sequencing technology, high-dimensional datasets, large-scale gene expression is accessible, which allowing us to detect aberrantly expressed MRGzs related to occurrence or metastasis of cancers and patients’ survival probability and the response to immunotherapy.

In the present study, we performed a comprehensive analysis in multiple databases and explored
the effect of MRGs on the survival of UM patients. Notably, the least absolute shrinkage and selection
operator (LASSO) Cox regression algorithm was used to analyze high-dimensional data. Then, a five-
gene prognostic model was constructed to generate a prognostics risk score, and patients were stratified
based on risk score, which was further validated in the gene expression omnibus (GEO) databases. In
addition, Cox regression analysis was used to relationship between the predictive value and clinical
information of UM patients. The nomogram was built for predicting the overall survival (OS) of UM.
Moreover, GSEA and immune infiltration analysis were used to identify the role of a signature in the
tumor microenvironment. The high-risk group patients have higher expression of immune checkpoint
molecules. In summary, this study provided a reliable five-genes signature based on MRGs that could
function as an independent prognostic marker for UM.

2. Materials and methods

2.1. Data collection from the cancer genome atlas (TCGA) and gene expression omnibus (GEO)
databases

The gene expression profiles and the corresponding clinical data of 80 UM patients were
downloaded from The University of California Santa Cruz (UCSC) Xena browser (https://xenabrowser.net/) with cohort name: TCGA-UVM. Then, the gene expression profiles of the
TCGA-UVM dataset (FPKM value) were transformed into transcripts per kilobase million (TPM)
value. Ensemble IDs converted to gene symbols using the “org.Hs.eg.db” and “clusterProfiler” R
packages. Besides, the gene expression profiles and survival data of GSE84976 (platform GPL10558,
Illumina HumanHT-12 V4.0 expression beadchip Illumina, Inc., San Diego, CA, United States) were
downloaded from GEO (https://www.ncbi.nlm.nih.gov/geo/) databases, which was chosen for this
study as the validation group. Strawberry Perl (version 5.32.0; http://strawberryperl.com/) was used to
extract the gene expression data from the TCGA-UVM and GSE84976 datasets and construct a data
matrix for further analysis. The MRGs were downloaded from the HALLMARK gene sets in the
Molecular Signature Database (MSigDB) (http://www.gsea-msigdb.org/gsea/msigdb/). Finally, a total
of 944 MRGs were identified for our study.

2.2. Protein-protein network interaction

The STRING (search tool for the retrieval of interacting genes/proteins) database (https://string-
db.org/) contains known and predicted PPI [15]. After constructing the PPI, the core genes of the
network need to be identified. We obtained the top 50 hub genes with the largest numbers of adjacent
nodes for subsequent analysis using R software.

2.3. Gene ontology (GO) and the Kyoto encyclopedia of genes and genomes (KEGG) pathway
functional enrichment analysis of hub genes

GO and KEGG enrichment analysis was performed utilizing hub genes. The KEGG results were
analyzed and visualized using the “ggplot2” and “clusterProfiler” R packages. The GO biological
process (BP), cellular component (CC), and molecular function (MF) results were visualized using the
“cnetplots” R package. We used p-value < 0.05 and FDR adjusted p-value < 0.05 as the GO and KEGG
enrichment analysis threshold.

2.4. Construction of the metabolism-related prognostic signature

The univariate Cox analysis was performed to screen out potential prognostics MRGs (p < 0.001) on the training cohort. Only genes that showed significantly associated with OS of patients with UM were considered for the subsequent analysis. Then, the LASSO Cox regression was used to screen the narrow candidate genes and avoid overfitting by using “glmnet” and “survival” R packages [16]. The penalty parameter lambda was detected by using 10-fold cross-validation [17]. The minimum lambda was defined as the optimal value, and we obtain a list of prognostic signatures with correlation coefficients. Next, the multivariate Cox regression analysis was used to identify the independent prognostic MRGs, and a prognostic model was constructed. The risk score of each patient was calculated as the following equation:

\[
\text{Risk score (patients)} = \sum_{i=1}^{n} \text{Expression}_{genei} \times \text{Coefficient}_{genei}
\]

Here, “\(n\)” represents the number of prognostic genes; “\(I\)” the serial number of each gene. UM patients were divided into high-risk and low-risk groups with the median risk score as a cutoff value according to the risk score formula. The Kaplan-Meier (K-M) survival curve was used to compare the prognostic gene signature and OS based on the two groups through the “survival” R package. Besides, the time-dependent receiver operating characteristic (ROC) curve was constructed by the “timeROC” R package, and the area under the curve (AUC) was calculated to measure the sensitivity and specificity of the multi-gene prognostic model.

2.5. Validation of the metabolism-related prognostic signature

To ensure the stability and repeatability of the multi-gene prognostic model, we calculated the risk score of each patient in GSE84976 cohorts, and patients were stratified into the high-risk and low-risk groups with the median risk score. The K-M curve and log-rank test were performed to evaluate the differences in survival rate between the two groups. The ROC curve and AUC analysis were implemented to show the predictive ability of prognostic signatures in the validation cohorts. The 1-, 3-, and 5-year survival rates of patients in the validation cohort were evaluated.

2.6. Gene set enrichment analysis (GSEA) analysis

GSEA is a computational method determining whether an a priori defined set of genes shows statistically significant, concordant differences between two biological states. The GSEA analysis was performed to identify differences in the set of genes expression between the high-risk and low-risk groups. In this study, the KEGG gene sets (v7.4) and HALLMARK gene sets (v7.4) were downloaded from the MSigDB. According to the GSEA user guide, NOM p-val < 0.05 and FDR q-val < 0.25 were considered as statistically significant. The top 5 pathways were selected and visualized.
2.7. Construction of the nomogram

In order to predict the 1-, 3-, and 5-year survival probability of UM patients, the nomogram was constructed based on the results of multivariate Cox regression analysis. Moreover, the calibration curve for 1-, 3-, and 5-year survival rates was used to access the prognostic accuracy of the nomogram. The nomogram and calibration curve were constructed by using “rms” R package.

Figure 1. Screening for metabolism-related genes and gene functional enrichment analysis. (A) protein-protein interaction of metabolism-related genes. (B) The top 50 core genes are based on the number of adjacent nodes for each protein. (C) The top 30 significant terms of GO function enrichment. BP, biological process; CC, cellular component; MF, molecular function. (D) The top 10 significant terms of KEGG analysis. KEGG, Kyoto Encyclopedia of Genes and Genomes.
2.8. Statistical analysis

All analyses were performed with R software (version 4.0.4, 64-bit; https://www.r-project.org/) and its appropriate packages. In addition to noting before, “ggplot2”, “ggpubr”, “limma”, “tidyverse”, “dplyr” and “plyr” R packages were also used for data analysis and graph plotting. The Perl programming language (version 5.34.0, https://www.perl.org/) was used to process data. The K-M curve and log-rank test were used to evaluate the differences in OS between the high-risk and low-risk groups. The Wilcoxon rank-sum test was used to verify the differences in expression of immune-related genes between the high-risk and low-risk groups. The p < 0.05 was considered statistically significant.

3. Results

3.1. Extraction and screening of metabolism-related genes

We first downloaded the KEGG metabolism-related genes from the MSigDB. A total of 944 MRGs were obtained from this gene set. Then, we used the STRING website to perform a PPI network analysis between MRGs (Figure 1A). After identifying the core genes of the network and the number of adjacent nodes for each protein, we obtained the top 50 core genes (Figure 1B). To understand the potential biological functions of these core genes, GO and KEGG analyses were conducted. The GO analysis showed that MRGs were closely involved in a variety of terms, including “nucleobase-containing small-molecule biosynthetic process” in the BP category, “mitochondrial matrix” in the CC category, and “coenzyme binding” in the MF category (Figure 1C). In the KEGG pathway enrichment analysis, MRGs are involved in material synthesis and material metabolism, including the “purine metabolism”, “carbon metabolism”, “pyrimidine metabolism” and et al. (Figure 1D).

3.2. Establishment of prognostic signature from the training cohort

We obtained the gene expression data and corresponding clinical information from the TCGA-UVM cohort. The univariate Cox proportional hazard regression analysis was performed to identify the association of MRGs with OS in UM patients. Our results showed that 15 MRGs were significantly associated with the OS of UM patients (Figure 2A). Among the survival-related genes, overexpression of 13 genes (OGHD, MDH1, TPI1, MDH2, HK1, NME3, NME1, ITPA, TXNRD1, GOT1, SHMT2, NT5C2, and GLUD1) was found to be significantly related to worse survival outcomes. In comparison, overexpression of 2 genes (PC and ENPP1) showed the prognostic value of indicating better OS. Then, we performed a LASSO regression analysis of these genes, and regression coefficients were calculated (Figure 2B). The LASSO analysis indicated that the model achieves the best performance when it includes 10 genes (Figure 2C). Finally, the multivariate Cox regression analysis was performed to construct the prognostic model, and five MRGs were identified as the independent prognostic factors (Figure 2D). The prognostic risk score was calculated as follow: Risk score = (expression level of MDH2 * 2.761058) + (expression level of NME1 * 1.384644) + (expression level of NT5C2 * 1.056515) + (expression level of PC * -1.443098) + (expression level of ENPP1 * -0.732329) (Table 1).
Table 1. Overall information of five metabolism-related genes is significantly associated with the prognosis of patients with uveal melanoma.

| Gene symbol | Description                                      | Ensembl ID       | Coefficient |
|------------|--------------------------------------------------|------------------|-------------|
| MDH2       | Malate dehydrogenase 2                           | ENSG00000146701  | 2.761058    |
| NME1       | NME/NM23 nucleoside diphosphate kinase 1         | ENSG00000239672  | 1.384644    |
| NT5C2      | Cytosolic 5’ nucleotidase II                     | ENSG00000076685  | 1.056515    |
| PC         | Pyruvate carboxylase                             | ENSG00000173599  | -1.443098   |
| ENPP1      | Ectonucleotide                                  | ENSG00000197594  | -0.732329   |

Figure 2. Establishment of metabolism-related prognostic signature from the training cohort. (A) Univariate Cox regression analysis identified candidate MRGs related to the UM risks. (B) Screening the Log Lambda value was corresponding to the minimum cross-validation error point. (C) Selecting the metabolic genes with a non-zero coefficient corresponding to the same Log Lambda value. (D) Multivariate Cox regression analysis revealed five independent MRGs related to patient prognosis. MRGs, metabolism-related genes; UM, uveal melanoma.
3.3. Effect of the metabolism-related signature on prognosis for UM

Figure 3. Construction and validation of a prognostic model related to OS based on metabolism-related genes. (A),(B) The Kaplan-Meier survival curves for patients with UM in the TCGA training cohort (A) and GEO validation database (B). (C),(D) The time-independent receiver operating characteristic curves of the prognostic signature in the TCGA training cohort (C) and GEO validation database (D). OS, overall survival; UM, uveal melanoma; TCGA, The Cancer Genome Atlas; GEO, Gene Expression Omnibus.

The five-genes prognostic signature model was established based on multivariate Cox regression analysis. According to the prognostic model, the risk score of each patient was calculated. The patient in TCGA and GEO cohorts were divided into high-risk and low-risk groups based on the median risk score value. To identify the survival difference between these two groups, we conduct a K-M curve analysis. Our results reveal significant differences between the high-risk and low-risk groups. Patients in the high-risk group show markedly poorer OS than those in the low-risk group (Figure 3A). Also, an unfavorable OS outcome was seen in the high-risk group of the testing group (Figure 3B).
Subsequently, the accuracy of the OS estimate derived from the prognostic model was performed through a time-dependent ROC curve. The ROC curve showed a gradually increasing accuracy of predicting the 1-year, 3-year, and 5-year survival rates. The AUC of the ROC curve was 0.861 at 1-year, 0.767 at 3-year, and 0.909 at 5-year in the training cohort (Figure 3C). The AUC of the ROC curve was 0.807 at 3-year and 0.861 at 5-year in the testing cohort (Figure 3D).

Furthermore, the distribution of survival status of the five-genes signature of patients both in training and testing cohort were plotted in Figure 4A,B. The histogram showed that a higher proportion of survival patients in the low-risk group than the high-risk group. We also analyzed the interaction between the MRGs identified as the prognostic model (Figure 4C,D). Then, the risk score, survival status, and gene expression were presented in Figure 4E and F. As showed in the figure, the survival time was longer in the low-risk group than in the high-risk group. In addition, the heatmap showed that NT5C2, MDH2, and NME1 were highly expressed in the high-risk group, PC and ENPP1 were highly expressed in the low-risk group.

3.4. Correlation of risk score with the clinical characteristics

The K-M survival analysis was applied to determine the relationship between OS rate in different subgroups of patients according to the risk score level and clinical characters, such as different stages, age, grade, gender, and TMN status. The results indicated that five-gene signature can significantly distinguish the prognosis of patients with the following characteristics: female, male, age \( \leq 65 \), age > 65, stage I-II, stage III-IV, T1-2, T3-4, M0, M1, N0, and N1, respectively (Figure 5A–L). Subsequently, the univariate and multivariate Cox regression analysis also indicated that the novel prognostic model could work as an independent prognostic factor related to the OS rate of UM patients (Figure 5M,N).

3.5. Correlation of risk score with immune checkpoint genes

Immunotherapy is a novel treatment for advanced tumors, activating the host’s natural immune defense system. However, the limitation of immunotherapy is that it can only benefit a minority part of patients. So, a more reliable biomarker to predict patient response to immunotherapy. Thus, we analyzed the expression level of the immune-related genes between high-risk and low-risk groups. A heatmap was then drawn to visualize the expression of these genes of patients in TCGA and GEO databases. As shown in Figure 6A, B, the expression level of EDNRB, MICA, and CCL28, which negatively regulated the trafficking and infiltration of immune cells into cancer, were significantly higher in high-risk groups than in low-risk groups group (\( p < 0.01 \)). Next, we investigated the expression of ICB molecules in the high-risk and low-risk groups. Our results showed that programmed cell death-1 (PD-1) was positively correlated with a risk score, was upregulated in the high-risk group (Figure 6C,D). In addition, the expression level of critical immune checkpoints (CD272, CD27, and IDO1) in the high-risk group was significantly higher than that in the low-risk group (Figures 6G–J). We also observed that immunosuppressive cytokines were also upregulated in the high-risk group (Figures 6K,L). Altogether, these results indicated that the patient with high-risk scores was found to tend to develop an immunosuppressive microenvironment through the upregulation of immunosuppressive cytokines and immune checkpoint-related genes.
Figure 4. Prognostic value of the metabolism-related risk signature in UM patients. (A),(B) Patient survival of the high and low hypoxia risk groups in the TCGA training cohort (A) and GEO validation database (B). (C),(D) Interaction analysis of the genes in the risk model on the TCGA training cohort (C) and GEO validation database (D). (E),(F) The distribution of risk score, survival status, and expression heatmap of the five MRGs in the TCGA training cohort (E) and GEO validation database (F). UM, uveal melanoma; TCGA, The Cancer Genome Atlas; GEO, Gene Expression Omnibus.
Figure 5. Relationship between the risk model and clinical factors. (A–L) The Kaplan–Meier analysis of clinical subgroup patients based on the risk model. The clinical characteristics including: female (A), male (B), age <= 65 (C), age > 65 (D), stage I-II (E), stage III-IV (F), T1-2 (G), T3-4 (H), M0 (I), M1 (J), N0 (K), and N1 (L). (M–N) Univariate and multivariate Cox prognostic analyses evaluating the independent risk factors of the metabolism-related signature in terms of OS in UM patients. OS, overall survival; UM, uveal melanoma.
Figure 6. Relationships between the risk score and the immune-related genes and immunosuppressive cytokines. (A–B) Heatmap of the expression levels of immune-related genes in different risk groups on the TCGA training cohort (A) and GEO validation database (B). (C–J) The correlation between the expression levels of immune checkpoint molecules and different risk groups on the TCGA training cohort and GEO validation database. (K–L) The immunosuppressive cytokine expression in the high-risk and the low-risk groups in the TCGA training cohort (K) and GEO validation database (L). TCGA, The Cancer Genome Atlas; GEO, Gene Expression Omnibus. *p < 0.05, **p < 0.01, and ***p < 0.001.
3.6. Gene set enrichment analysis

To understand the level of pathway enrichment between the high-risk and low-risk groups, the GSEA software was used to perform pathway enrichment analysis according to NES, nominal \( p \)-value, FDR \( q \)-value, and FWER \( p \)-value. As shown in Figure 7A, we found that all markedly enriched gene set of HALLMARK collection was seen in the high-risk group, such as angiogenesis, IL-6-JAK-STAT3 signaling, inflammatory response, oxidative phosphorylation, and UV response. The other related pathway annotated by the HALLMARK was shown in Supplementary Table S1. Moreover, all gene sets enriched in the KEGG collection were also enriched in the high-risk group (Figure 7B). These pathways mostly correlated with apoptosis, cell cycle, glutathione metabolism, mTOR signaling pathway, and p53 signaling pathway. The other related pathway annotated by the KEGG was shown in Supplementary Table S2,3.

3.7. Establishment and calibration of an integrated nomogram

Nomogram was constructed with the five prognostic genes to predict the OS rate of patients with UM. In the present study, a nomogram associated with the 1-year, 3-year, and 5-year survival rates was established (Figure 8A). Moreover, the calibration curve of the nomogram indicated an optimal agreement between the nomogram-predicted and observed OS rate (Figure 8B). The same results were seen in the validation cohort (Supplementary Figure S1).

**Figure 7.** GSEA enrichment analysis between low-risk and high-risk groups. (A) Five representative HALLMARK pathways in low-risk and high-risk patients. (B) Five representative KEGG pathways in low-risk and high-risk patients.
Figure 8. Development of a nomogram based on the metabolism-related signature in the TCGA training cohort. (A) Development of MRGs nomogram. (B–D) Calibration plots for the signature at 1-year, 3-year, and 5-year.

4. Discussion

UM is one of the highly aggressive tumors and remarkably variable in patient’s survival rate. The advancement of high-throughput technologies and bioinformatic methodology enables us to identify the crucial role of gene signatures based on specific associations in predicting the prognostic outcome of UM patients. Moreover, numerous HALLMARK gene sets related to tumor prognosis have been defined, such as autophagy, hypoxia, angiogenesis, and metabolism. In addition, the novel gene signature may also provide a more reliable and more effective biomarker in early detection and treatment.

Recently, the dysregulated metabolism of cancer has become a hotspot in cancer research, which might serve as a biomarker and a novel therapeutic strategy. Moreover, it has been reported to play a vital role in tumorigenesis, development, progression, and therapeutic resistance of different types of cancer. However, there is little known about the role of MRGs in the pathogenesis and progression of UM.

This study constructed a prognostic signature based on the MRGs in UM patients from the TCGA
and GEO databases. Using the univariate Cox regression analysis, we found 15 MRGs were significantly correlated with OS rate in UM patients. After conducting the LASSO regression and multivariate Cox regression analysis between the patient’s prognosis and gene expression, we obtained a novel prognostic signature, which consisted of five MRGs (NT5C2, MDH2, NME1, PC, and ENPP1). Then, we divided the UM cohort into the high-risk and low-risk groups according to the median risk score. The K-M survival curve analysis showed that the prognosis of patients in the high-risk and low-risk groups was significantly different. The OS rate in the low-risk group was markedly higher than that of their counterpart. Moreover, the time-dependent ROC curve, which aimed to evaluate the accuracy of the OS estimate, demonstrated that the AUC for 1-year, 3-year, and 5-year was 0.861, 0.767, and 0.909, respectively. Thus, the nomogram was constructed to represent the relations between MRGs expression level and prognosis. The calibration of the nomogram indicated that it has a good agreement between the predicted and observed survival rates. In addition, we performed a K-M analysis of the correlation of risk score with other clinical factors, such as gender, age, Stage, and TMN status. We also evaluated the prognostic value of this signature in the validation cohort, which indicated that the five-genes prognostic signature could be used as an independent prognosis indicator in both databases.

Five MRGs (NT5C2, PC, MDH2, ENPP1, and NME1) were selected as the prognostic signature in this study. Some of the genes in our signature have previously been shown to be involved in melanoma. However, several genes’ role in UM remains unclear. NT5C2, cytosolic 5’ nucleotidase II, plays an essential role in cellular nucleotide homeostasis. Recently studies revealed that NT5C2 is associated with cancer cell survival and chemotherapy resistance, including relapsed acute lymphoblastic leukemia and glioblastoma [18,19]. Pyruvate carboxylase (PC), a mitochondrial enzyme during the tricarboxylic acid cycle, plays a crucial role in catalyzes the ATP-dependent carboxylation of pyruvate to oxaloacetate [20]. Overexpression of PC is related to cancer cell survival and proliferation in non-small-cell lung cancer [21]. The PC inhibitor phenylacetic acid could suppress vemurafenib-treated melanoma cells growth, sensitizing melanoma cells to vemurafenib treatment [22,23]. Malate dehydrogenase 2 (MDH2), a member of malate dehydrogenase, is a crucial enzyme involved in the citric acid cycle in mitochondria [24]. Zhuang et al. reported that MDH2 could promote the proliferation, migration, and invasion but inhibited the apoptosis of the endometrial cancer cell line by suppressing PTEN [25]. The previous study also indicated that MDH2 is considered a candidate therapeutic target for cancer metabolism and tumor growth [26]. Ectonucleotide pyrophosphatase/phosphodiesterase 1 (ENPP1) is a type II transmembrane glycoprotein involved in insulin resistance and bone metabolism [27]. Abnormal expression of ENPP1 can increase human lung cancer malignancy via epithelial-mesenchymal transition [28]. In addition, the expression level of ENPP1 from patients with triple-negative breast cancer was significantly associated with recurrence-free survival and overall survival [29]. NME1 is a metastasis suppressor gene that inhibits tumor cells’ metastatic activity in different types of cancer. NME1 was associated with a statistically significant risk of breast cancer relapse and tumor metastasis [30]. NME1 was also able to suppress metastasis in a mouse model of UV-induced melanoma [31]. However, Wang et al. reported that NME1 could promote the expansion of melanoma cells, thus contributing to the melanoma growth and lung colonizing activities, which indicated NME1 could act as a driver of melanoma growth distinct from its function as a metastasis suppressor [32].

The clinical trial indicated that immunotherapy has high efficacy in tumor treatment and can improve the quality of life in patients with advanced melanoma. However, the main disadvantage of
immunotherapy is that only a minority of patients respond to it. Previous studies reported that immune checkpoint molecules could affect the metabolic communication of T cells. For instance, PD-1–PD-L1 axis might impair the metabolism pathway, such as aerobic glycolysis and glutaminolysis, in T cells through the PI3K–AKT–mTOR pathway [33,34]. In addition, to impair the metabolic profile of tumor-infiltrating lymphocytes, immune checkpoint molecules can also directly support T cell activation via reprogramming of metabolism. For example, CD28 signaling can enhance aerobic glycolysis and stimulate mitochondrial fusion of T cells [35]. In this study, we identified the association between risk score and immune checkpoint molecules. The high-risk group patients have a higher expression level of immune checkpoint molecules than those in the low-risk group.

Furthermore, we found different gene expression levels, which encode chemokines between the high-risk and low-risk groups. We found that the high-risk group patients have a high expression of six chemokines (CXCL9, CXCL10, CXCL11, CCR5, CCL20, CXCR3) low-risk group. Chemokines promote the migration, localization, and progression of several cancers. It has been reported that CCL20 is a poor prognostic factor for cutaneous melanoma [36]. CXCL10 is secreted by the various immune cell, is associated with melanoma invasion, proliferation, and metastasis [37]. It has also been observed that anti-PD-1 therapy is not effective without CXCL9, CXCL10, CXCL11/CXCR3 axis [38]. For anti-CTLA4 cancer immunotherapy, the expression levels of this axis were significantly increased in melanoma patients with good clinical responses [39]. CCR5 acts as a modulator of the immune response of interleukin-2 signaling therapy, contribute to the harmful inflammatory response [40]. Taken together, the outcome of our study revealed that the five-genes prognostic signature might be a potential predictive indicator in accessing the response of UM patients to immunotherapy.

However, there are some limitations to our study. First, the size of TCGA-UVM and GSE84976 cohorts is relatively small, which may not be consistent with the clinical population. Secondly, due to the limited clinical characteristic of patients in the GSE84976 cohort, some clinical subgroups analysis could not be performed. Finally, long-term prospective clinical research is needed to validate the robustness of the prognostic signature.

5. Conclusions

In summary, we constructed a novel five-gene signature prognostic risk model based on MRGs in the TCGA database that was an independent prognostic factor for UM patients and validated by an independent cohort from the GEO database. Our study may provide guidance for targeted therapy and potential biomarkers in the future.

Acknowledgments

This study was supported by the National Natural Science Foundation of China (Grant nos.82070924) and the Open Sharing Fund for the Large-scale Instruments and Equipment of Central South University.

Conflict of interest

The authors declare no competing interests.
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