The role of costimulatory molecules in glioma biology and immune microenvironment

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Background: Extensive research showed costimulatory molecules regulate tumor progression. Nevertheless, a small amount of literature has concentrated on the potential prognostic and therapeutic effects of costimulatory molecules in patients with glioma.

Methods: The data were downloaded from The Cancer Genome Atlas (TCGA) database, Chinese Glioma Genome Atlas (CGGA) database, and Gene Expression Omnibus (GEO) database for bioinformatics analysis. R software was applied for statistical analysis. Using the FigureYa and Xiantao online tools (https://www.xiantao.love/) for mapping.

Results: The Least absolute shrinkage and selection operator (LASSO) and Cox regression analysis were utilized to identify the signature consisting of five costimulatory molecules. Multivariate regression analysis revealed that the prognosis of glioma could be independently predicted by the riskscore. Furthermore, we explored clinical and genomic feature differences between the two groups. The level of tumor mutational burden (TMB) was higher in the high-risk group, while more mutation of IDH1 was observed in the low-risk group. Results of Tumor Immune Dysfunction and Exclusion (TIDE) analysis showed that high-risk patients were more prone to be responded to immunotherapy. In addition, subclass mapping analysis was performed to validate our findings and the results revealed that a significantly higher percentage of immunotherapy response rate was observed in the high-risk group.

Conclusion: A novel signature with a good independent predictive capacity of prognosis was successfully identified. And our findings reveal that patients with high-risk scores were more likely to be responded to immunotherapy.

Abbreviations: TCGA, The Cancer Genome Atlas database; CGGA, Chinese Glioma Genome Atlas; TMB, Tumor mutational burden; TIDE, Tumor Immune Dysfunction and Exclusion; WHO, World Health Organization; HGGs, High-grade gliomas; GBM, Glioblastoma; BBB, Blood-brain barrier; TNF, Tumor necrosis factor; ICI, Immune checkpoint inhibitors; PD-1, Programmed cell death protein 1; PD-L1, Programmed cell death 1 ligand 1; GTEx, Genotype-Tissue Expression; ssGSEA, Single-sample GSEA analysis; KM, Kaplan–Meier; ROC, Receiver operating characteristics; AUC, Area under Gene set enrichment analysis; LASSO, Least absolute shrinkage and selection operator.
Introduction

Gliomas are the most common and aggressive primary tumors of brain, with approximately 200,000 deaths worldwide (Sung et al., 2021). Generally, according to the World Health Organization (WHO) standards, grades I-II are low-grade gliomas (LGGs) and grades III-IV are high-grade gliomas (HGGs) (Figarella-Branger et al., 2021). It is worth noting that glioblastoma (GBM) with WHO grade IV, well known for their aggressiveness and high propensity to metastasize, have only a 5% five-year survival rate (Shergalis et al., 2018). Despite major advances in cancer treatment and many emerging therapies that have been proposed for glioma patients, the overall survival rate (OS) has not significantly increased in recent years (Xu et al., 2020). Under most circumstances, patients with glioma have advanced into the intermediate and advanced stages at the time of diagnosis, missing out on the optimal treatment time (Wang et al., 2020a). Currently, WHO grade is most commonly used as a reference to judge the clinical prognosis of glioma patients, which is valuable but insufficient for prognostic prediction and evaluation of subgroups of patients (Ali et al., 2021). Thus, it is important to find additional more effective targets and more susceptible therapeutic options for glioma patients to improve the prognosis.

With the development of molecular biology, immunotherapy has been recognized as a promising treatment for overcoming glioma despite the presence of the blood-brain barrier (BBB) (Ye et al., 2019; Marcucci et al., 2021). Costimulatory molecules and signals consisting of the tumor necrosis factor (TNF) families and B7/CD28 family are promising candidates for immunotherapy (Croft et al., 2013). On the one hand, molecules belonging to B7/CD28 family are essential for triggering immune responses including the most common immune checkpoint inhibitors (ICIs) target PD-1 and PD-L1 (Keir et al., 2008). In another hand, molecules belonging to TNF/TNFR family are essential for the promotion of anti-tumor immunity (Driessens et al., 2009). In recent years, many literatures show that costimulatory molecules are linked to the tumorigenesis and promotion of various cancers. CD40LG (CD40L, TNFSF5, CD154), one of the most well-studied TNFSF, has been a therapy target in cancer treatment and is typically associated with the prognosis of lung cancer (Mu et al., 2015). Overexpression of TNFSF14 can stop or delay the development of human papillomavirus 16-induced tumors via enhancing the functional responses of T cells (Kanodia et al., 2010). Moreover, EDAR is an important effector of typical Wnt signaling in the development of skin attachment, which can adjust Wnt/β-catenin signaling pathway to promote the proliferation of colorectal cancer cells (Wang et al., 2020b). Given the prominent values of costimulatory molecules, it is essential to screen the costimulatory molecules associated with the prognosis for improving prognosis evaluations of glioma patients.

To investigate the significant role of costimulatory molecules in glioma, RNA sequencing data was used to systematically analyze the costimulatory molecules expression with distinct clinicopathological features of gliomas in four independent cohorts according to the TCGA and CGGA datasets. A prognostic signature was then developed that could effectively predict the glioma prognosis. Moreover, we investigated discrepancies in clinical and genomic profiles in two risk groups and explored potential targets for therapies.

Materials and methods

Data acquisition and processing

The relevant clinical data and RNA-sequencing data of TCGA-LGG patients and TCGA-GBM patients were acquired from the TCGA database (https://portal.gdc.cancer.gov/) as the training cohort. In addition, four datasets including data of glioma were retrieved from the CGGA and GEO databases (https://cgga.org.cn/ and https://www.ncbi.nlm.nih.gov/geo/) as the validation cohorts. The Genotype-Tissue Expression (GTEx) database (https://gtexportal.org/) RNA-Seq data was also downloaded for further analysis. Then, data was merged, converted to TPM values (Zhao et al., 2020), annotated with probes, and removed samples with incomplete information and duplicates. The data were then batch normalized using the “ComBat” algorithm to decrease the possibility of batch effects in disparate datasets. The expression profile of patients who responded to immunotherapies was collected from the known literature and was applied for predicting the immunotherapy response of glioma patients (Roh et al., 2017). Single-sample GSEA analysis (ssGSEA) was conducted via the “GSVA” R package (version 4.0.2).

Differential expression analysis

All costimulatory molecules were gotten from the published literature (Aye et al., 2021). RNA sequencing expression data of 56 costimulatory molecules in our research including normal samples and tumor samples were obtained via UCSC XENA data hubs (https://tcga.xenahubs.net). Then differential expression analysis of all these costimulatory molecules was conducted via the “EdgeR” R package (version 4.0.2), and we visualized the results via the “ggplot2” R package (version 4.0.2).
Establishment and evaluation of the prognostic feature in a combined glioma cohort

After eliminating patients with survival time less than 30 days and deficient clinic information, 622 patients in the TCGA database were used to establish a prognostic signature, and 929 patients in the CGGA database were utilized to estimate the predictive ability and reliability of the signature. First, univariate Cox regression was conducted to obtain the costimulatory molecules related to OS of glioma patients \( (p < 0.01) \). The LASSO regression analysis was then conducted via the “glmnet” R package (version 4.0.2) to confirm the selected hub genes further. Finally, the prognostic signature was defined via a multivariate Cox regression analysis and the riskscore was obtained for each patient using the following formula: 

\[
\text{Riskscore} = \text{coef1} \times \text{costimulatory molecule} + \text{coef2} \times \text{costimulatory molecule2} + \text{coef3} \times \text{costimulatory molecule3} + \ldots + \text{coefN} \times \text{costimulatory molecules}.
\]

Differentially expressed genes with \( p \)-value < 0.05 in two risk groups were screened. Differential expression analyses of all samples, and 1,152 normal samples from the GTEx database, 689 tumor samples from TCGA and CGGA databases. Meanwhile, we found an evident batch effect in the four datasets (Figure 1A). The “sva” R package was utilized for eliminating the potential batch effect, and inter-assay differences were found significantly decreased in the conjunct glioma cohort (Figure 1B). Moreover, the expression of costimulatory molecules in each patient was quantified by the ssGSEA algorithm for further analysis (Figure 1C).

GSEA and mutation analysis

The “ClusterProfiler” R package (version 4.0.2) was utilized for performing GSEA. Curated gene sets, oncogenic signature gene sets, ontology gene sets and hallmark gene set (https://www.gsea-msigdb.org/gsea/downloads.jsp) was recognized as the reference sets to investigate the discrepancy in the tumor genetic pathways. Immunotherapy-related positive signatures from the known literature were obtained to conduct the correlation analysis with riskscore (Hu et al., 2021). Moreover, somatic mutation data used to calculate tumor mutational burden (TMB) were available from the cBioPortal website (https://www.cbioportal.org/datasets). Differentially mutated genes with \( p \)-value < 0.05 in two risk groups were screened and maftools were applied for analyzing the interaction between gene mutations.

Investigation of immune signatures and immunotherapeutic response prediction

The scores of ESTIMATE, immune, and stromal were counted by “estimate” R package (version 4.0.2). The ssGSEA algorithm was applied for quantifying the enrichment scores of 39 immune signatures. The 48 immune-checkpoint-relevant genes expression were selected for disparate expression analysis in two risk groups. TIDE analysis could precisely model immune escape and predict cancer response to immunotherapy (Jiang et al., 2018). Patients with TIDE score > 0 were considered to have no immunotherapy response, and patients with TIDE score < 0 were considered to have the immunotherapy response. Moreover, a subclass mapping algorithm (https://cloud.genepattern.org/gp), was utilized for determining which group was more likely to benefit from immunotherapy (Hoshida et al., 2007). In addition, appropriate targeted drugs were defined by differential expression analysis of drug sensitivity.

Statistical analysis

According to the R software (version 4.0.2), all analyses were conducted. All statistical tests were two-sided, and the difference was statistically significant when \( p \)-value < 0.05. Continuous variables with normal distribution were contrasted via an independent \( t \)-test. Wilcoxon rank-sum test was applied for comparing continuous variables.

Results

Data combination and correction for batch effect

After comprehensively retrieving the TCGA and CGGA databases, four glioma cohorts conformed to our standard, including TCGA-GBM, TCGA-LGG, CGGA_325, and CGGA_693 databases. Meanwhile, we found an evident batch effect in the four datasets (Figure 1A). The “sva” R package was applied for eliminating the potential batch effect, and inter-assay differences were found significantly decreased in the conjunct glioma cohort (Figure 1B). Moreover, the expression of costimulatory molecules in each patient was quantified via the ssGSEA algorithm for further analysis (Figure 1C).

Costimulatory molecules-based prognosis signature

After a comprehensive search performed on public databases, 1,152 normal samples from the GTEx database, 689 tumor samples, and five adjacent samples from the TCGA database were screened. Differential expression analyses of all costimulatory molecules were performed between normal samples from TCGA and GTEx databases, and tumor samples from the TCGA database. 57 differentially expressed costimulatory molecules with \( p < 0.001 \) were identified.
among which the expression of six costimulatory molecules including HHLA2, TNFRSF14, TNFRSF18, TNFRSF25, TNFRSF6B, and TNFSF9 were decreased in tumor samples (Figure 2A), whereas other costimulatory molecules were increased in tumor samples (Figures 2B–E). Patients with intact survival information were screened for further analysis (Supplementary Table S1). The training cohort was the TCGA cohort and the verification cohort was the CGGA cohort. Univariate Cox regression analysis was performed on the differentially expressed costimulatory molecules and
31 costimulatory molecules were screened (Figure 3A). Next, 11 OS-related costimulatory molecules were identified by LASSO Cox regression analysis (Figures 3B, C). According to five costimulatory molecules defined by multivariate Cox regression analysis, a prognosis signature was constructed and the risk score was calculated by the formula: risk score = CD274 * 0.198666431 + TNFRSF11B * 0.207922941 + TNFRSF14 * 0.18558208 + TNFRSF19 * 0.130431237404669 + TNFRSF21 * −0.1078322 (Figure 3D). Then, KM survival analysis of these five costimulatory molecules was conducted to contrast the OS time between two groups (Figure 3E).

**Validation of the prognostic feature**

The optimal truncation value of risk score 1.07792 in the training cohort and 0.159026 in the validation cohort were counted via the “maxstat” R package. Depended on the
optimal truncation value, patients were classified into high- and low-risk groups. Clinical information analysis indicated that high-risk patients may have adverse events (Figure 4A). KM survival analysis suggested that in the training cohort (Figure 4B) and validation cohort (Figure 4C), the OS time was markedly shorter in the high-risk group, showing that riskscore could predict the prognosis. As the riskscore increased, there was also a prominent increase in mortality in both the training cohort (Figure 4D) and validation cohort (Figure 4E). Consistently, the AUC value of 1-, 3-, and 5 years were respectively 0.70, 0.76, and 0.75 in the training cohort (Figure 4F) and 0.73, 0.77, and 0.79 in the validation cohort (Figure 4G). Moreover, to verify results in microarray datasets, GSE4290 and GSE108474 series were selected to perform the differential expression analysis and the clinical analysis, from which we found the expression of the signature costimulatory molecules was significantly different between tumor and normal tissues (Supplementary Figures S1A,B), and patients with advanced gliomas have a higher riskscore (Supplementary Figures S1C,D). Furthermore, given that the difference between two risk groups may just cause by natural difference between HGG and LGG, we conducted KM and ROC analysis in GBM and LGG patients independently, which were matching our results (Supplementary Figures S2A–F). Univariate and multivariate Cox regression analyses were conducted to demonstrate the possibility of the riskscore being a prognostic element distinct from common clinical factors. With the lowest p-value both in univariate and multivariate Cox regression analyses, the riskscore was linked to the prognosis (Figures 4H,I).

Enrichment analysis and tumor mutation burden

The enrichment scores of known immunotherapy-related positive signatures were calculated for each patient and we found
that riskscore positively correlated with all these signatures (Figure 5A), indicating that high-risk patients could be more suitable for anti-tumor immunotherapy. Furthermore, we also analyzed the correlations between riskscore and the scores of the hallmark gene set (Figure 5A). To better understand the biological processes and pathways of riskscore on the patient prognosis in two risk groups, oncogenic signature gene sets, ontology gene sets and curated gene sets were selected to perform GSEA analysis both in high- and low-risk groups patients (Figures 5B–G). Figure 6A showed the TMB level of all cancer types in the TCGA database, from which a clear difference in TMB value was observed between LGG and GBM patients, and
the TMB level in high-risk patients was evidently higher than that in low-risk patients (Figure 6B). The somatic mutation profiles revealed that IDH1 mutations were more frequent in low-risk patients and high-risk patients possessed specific top mutated genes (Figure 6C). Given the difference in TMB value between LGG and GBM patients, we then performed differential mutation analysis between two risk groups both in LGG and GBM patients. IDH1, EGFR, CIC, NF1, ATRX, TP53, FUBP1, and SMARCA4 in LGG patients, and ATRX, IDH1, UGGT1, DNAH7, ROBO1, PCNT, EP400, DSG3, SLIT3, RB1, VWF, TAF1L, and HMCN1 in GBM patients were identified that mutated significantly different between two risk groups (Figures 6D,E). Moreover, co-occurrences were confirmed in the mutations of these genes (Figures 6F,G).
Immunity and immunotherapy response analysis

The ESTIMATE algorithm was firstly conducted to quantify the immune scores, ESTIMATE scores, and stromal scores, and all the scores were higher in high-risk score patients (Figure 7A). Then, according to the ssGSEA Z-scores of 39 immune signatures, glioma patients were segmented into high- and low-immunity groups. We could see that high-risk patients consisted of more proportions of the high-immunity group (Figure 7B). Between the two risk groups, there were evident disparities in 39 immune signatures including immune functions and immune cells (Figures 7C,D). In light of the significant role of ICIs therapy in cancer, the expression distribution of 48 immune-checkpoint-correlated genes was presented in Figure 7E. Further, the correlation analysis showed that riskscore had a strong association with PD-1, PD-L1, PD-L2, and CTLA4 expression (Figure 7F). Most notably, all these findings revealed that high-risk score patients had a stronger immune-signature infiltration. Figure 8A showed the distribution of the TIDE score. In the high-risk group, there were 71.01% of patients responded to immunotherapy, while in the low-risk group, only 53.96% of patients responded to immunotherapy (Figure 8B). Following,
we conducted a TIDE analysis of our combined cohort, and the result revealed that the high-risk score linked to the low-TIDE score, demonstrating that these patients may be more likely to respond to immunotherapy (Figure 8C). Moreover, the TIDE analysis in GBM and LGG patients showed the same results independently (Supplementary Figures S3A,B). Besides the TIDE prediction, subclass mapping analysis was applied for verifying our findings, and we were delighted to see that high-risk patients were more potential to profit from PD-1 checkpoint treatment both in LGG and GBM patients (Figures 8D,E). Finally, the IC50 for each sample of 179 drugs were estimated via R "oncoPredict" package (version 4.0.2) in the GDSC database and between the two risk groups, drugs that have disparities in sensitivity were identified. The drugs with the most prominent sensitivity discrepancies in the two risk groups were displayed in Figures 8F,G.
Immunotherapeutic response prediction and screening of potential drugs. (A) Distribution of TIDE score. (B) Patients in the high-risk group have a higher percentage of responders. (C) Patients in the high-risk group have a lower level of TIDE score. (D,E) Subclass mapping analysis indicated patients in the high-risk group could be more sensitive to the PD-1 inhibitor in both LGG and GBM. (F,G) Drugs with the most significant sensitivity differences in high- and low-risk groups.
Discussion

With the most pernicious kind of primary brain cancer, glioma is famous for its high possibility of metastasis and recurrence (Yang et al., 2022). In the one hand, most glioma patients have developed to the intermediate and advanced stages at the time of diagnosis, missing the optimal treatment time. In another hand, glioma patients lack targeted therapy for specific subtypes and the current treatment can not control glioma invasion to normal adjacent brain tissue (Lee et al., 2018). The prognosis for patients with glioma remains dismal due to intratumoral heterogeneity (Jan et al., 2010). Accumulating evidence revealed a prominent association between different molecular subtypes and clinical outcomes in glioma patients (Li et al., 2022; Zheng et al., 2022). And several studies showed that costimulatory molecules took a great part in initiating anti-tumor immune responses (Zhang et al., 2020), but there was little literature focused on the correlation between costimulatory molecules and stratification of patients with glioma. This study discovered a novel signature that could effectively reflect the survival and immunotherapy response of glioma patients.

In our study, we firstly combined four independent cohorts into a large glioma cohort and used “sva” R package to decrease bias resulting in the small sample size. Univariate Cox analysis verified 31 costimulatory molecules closely correlated to patients’ prognosis and five costimulatory molecules including CD274, TNFRSF11B, TNFRSF14, TNFRSF19, and TNFRSF21 were selected by multivariate Cox analysis for signature construction. Recent research revealed that the expression of CD274 induced under hypoxia condition was signal associated with poor survival in glioma patients (Ding et al., 2021). In addition, TNFRSF11B might be involved in the malignant progression of gliomas and was one of the signature genes that predicted patient prognosis (Kang et al., 2021). TNFRSF14 served a tumor suppressive role by suppressing tumor cell proliferation and inducing apoptosis in bladder cancer and could act as a new diagnosis and prognostic biomarker for bladder cancer (Zhu and Lu, 2018). Further, TNFRSF19, essential for cell proliferation and development of nasopharyngeal carcinoma, represented a mechanism for tumor cells to escape from TGF-β growth-inhibitory action (Deng et al., 2018). Besides, a study demonstrated that the TNFRSF21 expression was strongly negatively linked to the miR-20a-5p expression, and the downregulation of TNFRSF21 functioned as an oncogene in squamous cell carcinoma of the head and neck (Wu et al., 2018). With the in-depth study of costimulatory molecules, the significant costimulatory molecules verified in this study might provide a foundation for further exploring the prognostic and therapeutic role of glioma.

We further investigated the potential biological discrepancy between the two risk groups through GSEA and mutation burden analysis. We found that riskscore positively associated with all known immunotherapy correlated positive signals. With the curated gene sets being a reference set, GSEA analysis indicated that

Glioblastoma_Mesenchymal, Nakayama_Soft_Tissue_Tumors, PCA1_Up, Noushmehr_GBM_Silenced_by_Methylayion, Blanco_Melo_COVID-19_Sars, Melachlan_Dental_Caries_Up pathways were evidently stimulated in high-risk patients. And Acute_Inflammatory_Response, B_cell_Receptor_Signaling_Pathyway, B_Cell_Mediated_immunity, Adaptive_Immune_Response_Based_On_Somatic_Recombination and Adaptive_Immune_Response were also significantly activated in high-risk patients with the ontology gene sets being a reference set. These findings might partially explain the worse prognosis of high-risk patients and suggested that these patients might be more suitable for anti-tumor immunotherapy. In addition, we found that the TMB level was evidently higher in high-risk patients. Besides, IDH1, associated with a good prognosis in glioma patients, was also observed in more mutations in low-risk patients (Bai et al., 2016).

Moreover, we explored the situation of immune cells and related immune pathways. Results suggested that high-riskscore patients got a higher level of most immune features. We also observed higher levels of PD-L1 and PD-L2 in high-risk patients. Studies had demonstrated that the PD-L1 was a predictive marker for tumor immunotherapy (Patel and Kurzrock, 2015). A study also reported that PD1 expression increased neuronal killing of cancer cells and was associated with long-term survival (Kingwell, 2013). The results of TIDE analysis suggested that in the high-risk group, patients had a higher immunotherapy response rate, which might be correlated to higher immune-checkpoint-related gene levels in high-risk patients. In addition, the subclass mapping algorithm analysis verified that patients with high-risk score were more potential to benefit from PD-1 checkpoint therapy.

To sum it up, this study successfully constructed and validated costimulatory molecules based on the prognostic feature, which might be applied for further guiding treatment and improving clinical outcomes for glioma patients. However, there are several limitations to the study. Firstly, because only individuals from Western and China populations are included, the samples may generate some population and genome bias. Then, this study lacks verification from other clinical data sets which will be beneficial to our signature. Finally, given the incompleteness of patient information and the sensitivity to incorrect model specification in our multivariate regression analysis, further analysis of patients with complete clinical information can be very beneficial in the future. Therefore, further investigation of prospective studies and other costimulatory molecules are needed such as functional experiments and underlying molecular mechanisms.

Conclusion

In conclusion, we investigated the biological features and prognostic value of costimulatory molecules in patients with glioma. We developed a new prognostic signature, and
demonstrated the potential immune-related mechanisms of this signature. Then, most importantly, our findings indicated that high-riskscore patients were more likely to benefit from immunotherapy.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

Author contributions

CT conceived the project. JW and ZW analyzed the data. Both JW, ZW, and WJ wrote and revised the article. WG, BD, ZW, and MZ performed the literature investigation and revised the article, including figures and tables. All authors have reviewed the article and all approved the final version.

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References

Ali, H., Harting, R., de Vries, R., Ali, M., Wurdinger, T., and Best, M. G. (2021). Blood-based biomarkers for glioma in the context of gliomagenesis: A systematic review. Front. Oncol. 11, 665235. doi:10.3389/fonc.2021.665235

Aye, L., Song, X., Yang, J., Hu, L., Sun, X., Zhou, J., et al. (2021). Identification of a costimulatory molecule gene signature to predict survival and immunotherapy response in head and neck squamous cell carcinoma. Front. Cell. Dev. Biol. 9, 695533. doi:10.3389/fcell.2021.695533

Bai, H., Harmananc, A., Erogn-Omay, E., Li, J., Cojkan, S., Simon, M., et al. (2016). Integrated genomic characterization of IDH1-mutant glioma malignant progression. Nat. Genet. 48 (1), 59–66. doi:10.1038/ng.3457

Croft, M., Benedict, C. A., and Ware, C. F. (2013). Clinical targeting of the TNF and TNFR superfamilies. Nat. Rev. Drug Discov. 12 (2), 147–168. doi:10.1038/nrd3930

Deng, C., Lin, Y., Qi, X., He, G., Zhang, Y., Zhang, H., et al. (2018). TNRFSF19 inhibits TGFβ signaling through interaction with TGFβ receptor type I to promote tumorigenesis. Cancer Res. 78 (13), 3469–3480. doi:10.1158/0008-5472.CAN-17-3205

Ding, X., Wang, L., Zhang, X., Xu, J., Li, P., Liang, H., et al. (2021). The relationship between expression of PD-L1 and HIF-1α in glioma cells under hypoxia. J. Hematol. Oncol. 14 (1), 92. doi:10.1186/s13045-021-01102-5

Driessens, G., Kline, J., and Gajewski, T. F. (2009). Costimulatory and coinhibitory receptors in anti-tumor immunity. Immunol. Rev. 229 (1), 126–144. doi:10.1111/j.1600-065X.2009.00771.x

Figarella-Branger, D., Appay, R., Metais, A., Tauriède-Espariat, A., Colin, C., Rousseau, A., et al. (2021). The 2021 WHO classification of tumours of the central nervous system. Ann. Pathol. 42 (5), 367–382. doi:10.1016/j.annpat.2021.11.005

Hoshida, Y., Brunet, J. P., Tamayo, P., Golub, T. R., and Mesirov, J. P. (2007). Subclass mapping: Identifying common subtypes in independent disease data sets. PLoS one 2 (11), e1195. doi:10.1371/journal.pone.0011195

Hu, J., Yu, A., Othmane, B., Qi, D., Li, H., Li, C., et al. (2021). Siglec15 shapes a non-inflamed tumor microenvironment and predicts the molecular subtype in bladder cancer. Theranostics 11 (7), 3089–3108. doi:10.7150/thno.53649

Jan, H. J., Lee, C. J., Shih, Y. L., Huang, D. Y., Ma, H. I., Lai, J. H., et al. (2010). Osteopontin regulates human glioma cell invasiveness and tumor growth in mice. Neuro. Oncol. 12 (1), 58–70. doi:10.1093/neuonc/nop013

Jiang, P., Yu, S., Pan, D., Fu, J., Sahu, A., Hu, X., et al. (2018). Signatures of T cell dysfunction and exclusion predict cancer immunotherapy response. Nat. Med. 24 (10), 1550–1558. doi:10.1038/nm.41591-018-0136-1

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2022.1024922/full#supplementary-material

SUPPLEMENTARY FIGURE S1

Differential expression analyses and the clinical analyses in microarray datasets. (A,B) Results of differential expression analyses showed that patients with glioma had a higher level of the signature genes in tumor tissues both in GSE4290 (A) and GSE108474 series (B). And patients with advanced stage qphad a higher level of riskscore (C,D).

SUPPLEMENTARY FIGURE S2

The validation of the model using LGG and GBM data independently. (A,B) KM survival curve analysis of the model. (C,D) The plots shows that the mortality rate of patients increases gradually as their riskscore increases. (E,F) 1, 3, and 5 years ROC curve analysis of the model in both LGGs and GBM data independently.

SUPPLEMENTARY FIGURE S3

The results of TIDE analyses in LGG and GBM patients independently. (A,B) The TIDE analyses showed that patients in high risk group with LGG or GBM all had a lower TIDE score.
Kang, K., Xie, F., Wu, Y., Wang, Z., Wang, L., Long, J., et al. (2021). Comprehensive exploration of tumor mutational burden and immune infiltration in diffuse glioma. *Int. Immunopharmacol.* 96, 107610. doi:10.1016/j.intimp.2021.107610

Kanodia, S., Da Silva, D. M., Karamanukyan, T., Bogaert, L., Fu, Y. X., and Kast, W. M. (2010). Expression of LIGHT/TNFSF14 combined with vaccination against human papillomavirus Type 16 E7 induces significant tumor regression. *Cancer Res.* 70 (10), 3955–3964. doi:10.1158/0008-5472.CAN-09-3737

Keir, M. E., Butte, M. J., Freeman, G. J., and Sharpe, A. H. (2008). PD-1 and its ligands in tolerance and immunity. *Annu. Rev. Immunol.* 26, 677–704. doi:10.1146/annurev.immunol.26.021607.090331

Kingwell, K. (2013). Neuro-oncology: Glioblastoma prognosis linked to neuronal PD-L1 expression in tumour-adjacent tissue. *Nat. Rev. Neurool.* 9 (11), 602–603. doi:10.1038/nrneurol.2013.197

Lee, E., Yong, R. L., Paddison, P., and Zhu, J. (2018). Comparison of glioblastoma (GBM) molecular classification methods. *Semin. Cancer Biol.* 53, 201–211. doi:10.1016/j.semcancer.2018.07.006

Li, X., Wang, Y., Wu, W., Xiang, J., Qi, L., Wang, N., et al. (2022). A novel risk score model based on eleven extracellular matrix related genes for predicting overall survival of glioma patients. *J. Oncol.* 2022, 4966820. doi:10.1155/2022/4966820

Marcucci, F., Corti, A., and Ferrari, A. J. M. (2021). Braching the blood-brain tumor barrier for tumor therapy. *Cancers* 13 (10), 2391. doi:10.3390/cancers13102391

Mu, C. Y., Qin, P. X., Qu, Q. X., Chen, C., and Huang, J. A. (2015). Soluble CD40 in plasma and malignant pleural effusion with non-small cell lung cancer: A potential marker of prognosis. *Exp. Ther. Med.* 9 (11), 602–603. doi:10.3892/etm.2020.9266

Patel, S., and Kurzrock, R. (2015). PD-L1 expression as a predictive biomarker in cancer immunotherapy. *Mol. Cancer Ther.* 14 (4), 847–856. doi:10.1158/1535-7163.MCT-14-0983

Roh, W., Chen, P. L., Reshef, A., Spencer, C. N., Prieto, P. A., Miller, J. P., et al. (2017). Integrated molecular analysis of tumor biopsies on sequential CTLA-4 and PD-1 blockade reveals markers of response and resistance. *Sci. Transl. Med.* 9 (379), eaaih3560. doi:10.1126/scitranslmed.aah3560

Shergalis, A., Bankhead, A., 3rd, Luesakul, U., Muangsin, N., and Neamati, N. (2018). Current challenges and opportunities in treating glioblastoma. *Pharmacol. Rev.* 70 (3), 412–445. doi:10.1124/pr.117.014944

Song, H., Ferluy, J., Siegel, R. L., Lavranos, M., Soerjomataram, I., Jemal, A., et al. (2021). Global cancer statistics 2020: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. *Ca. Cancer J. Clin.* 71 (3), 209–249. doi:10.3332/caac.21660

Wang, B., Liang, Y., Chai, X., Chen, S., Ye, Z., Li, R., et al. (2020). Ectodysplasin A receptor (EDAR) promotes colorectal cancer cell proliferation via regulation of the Wnt/β-catenin signaling pathway. *Exp. Cell. Res.* 395 (1), 112170. doi:10.1016/j.yexcr.2020.112170

Wang, Z., Wu, H., Yan, H., Cai, T., Dai, J., and Liu, Q. (2020). LINC00210 exerts oncogenic roles in glioma by sponging miR-328. *Exp. Ther. Med.* 20 (6), 137. doi:10.3892/etm.2020.9266

Wu, H., Pang, P., Liu, M., Wang, S., Jin, S., Liu, F., et al. (2018). Upregulated miR-20a-5p expression promotes proliferation and invasion of head and neck squamous cell carcinoma cells by targeting of TNFRSF21. *Onco. Rep.* 40 (2), 1138–1146. doi:10.3892/or.2018.6477

Xu, S., Tang, L., Ji, X., Fan, F., and Liu, Z. (2020). Immunotherapy for glioma: Current management and future application. *Cancer Lett.* 476, 1–12. doi:10.1016/j.canlet.2020.02.002

Yang, Z., Chen, Z., Wang, Y., Wang, Z., Zhang, D., Yue, X., et al. (2022). A novel defined pyroptosis-related gene signature for predicting prognosis and treatment of glioma. *Front. Oncol.* 12, 717926. doi:10.3389/fonc.2022.717926

Ye, L., Park, J. J., Dong, M. B., Yang, Q., Chew, R. D., Peng, L., et al. (2019). In vivo CRISPR screening in CD8 T cells with AAV-Sleeping Beauty hybrid vectors identifies membrane targets for improving immunotherapy for glioblastoma. *Nat. Biotechnol.* 37 (11), 1302–1313. doi:10.1038/s41587-019-0246-4

Zhang, C., Zhang, Z., Sun, N., Zhang, Z., Zhang, G., Wang, F., et al. (2020). Identification of a costimulatory molecule-based signature for predicting prognosis risk and immunotherapy response in patients with lung adenocarcinoma. *Oncoimmunology* 9 (1), 1824641. doi:10.1080/2162402X.2020.1824641

Zhao, S., Ye, Z., and Stanton, R. (2020). Misuse of RPKM or TPM normalization when comparing across samples and sequencing protocols. *RNA* 26 (8), 903–909. doi:10.1261/rna.074922.120

Zheng, J., Wu, Z., Qiu, Y., Wang, X., and Jiang, X. (2022). An integrative multiomics analysis based on liquid-liquid phase separation delineates distinct subtypes of lower-grade glioma and identifies a prognostic signature. *J. Transl. Med.* 20 (1), 55. doi:10.1186/s12967-022-03266-1

Zhu, Y., and Lu, M. (2018). Increased expression of TNFRSF14 indicates good tumor regression. *Mol. Cancer Ther.* 17 (1), 36–41. doi:10.1158/1535-7163.MCT-14-0983

Zhu, Y., and Lu, M. (2018). Increased expression of TNFRSF14 indicates good prognosis and inhibits bladder cancer proliferation by promoting apoptosis. *Mol. Med. Rep.* 18 (3), 3403–3410. doi:10.3892/mmr.2018.9306