Introduction: Glioblastoma (GBM) is a highly malignant primary brain tumor. Even after undergoing surgery and chemotherapy, patients with this affliction still have little to no chance of survival. Current research on immunotherapy treatment for GBM shows that immune-checkpoint inhibitors (ICIs) may be a promising new treatment method. However, at present, the relationship between the fatty acid metabolic process and the prognosis of GBM patients who are receiving immunotherapy is not clear.

Methods: First, we downloaded a GBM cohort that had been treated with immunotherapy, which included the mutation and prognosis data, and the TCGA-GBM and Jonsson-GBM queues. CIBERSORT and single sample gene set enrichment analysis (ssGSEA) were used to evaluate immune cell scores. Gene set enrichment analysis (GSEA) was used to evaluate the patient's accessment score. The pRRophetic algorithm was used to evaluate the drug sensitivity of each patient. Univariable and multivariate cox regression analyses, as well as the Kaplan-Meier (KM) method, were used to evaluate the relationship between the fatty acid metabolic process and the prognosis of GBM patients.

Results: The univariate and multivariate cox regression models showed that the fatty acid metabolic process mutant-type (MT) can be used as an independent predictor of the efficacy of immunotherapy for GBM patients. In addition, fatty acid metabolic process MT is related with significantly longer overall survival (OS) time than the wild-type (WT) variant. However, the mutation status of the fatty acid metabolic process has nothing to do with the prognosis of GBM patients who are receiving conventional treatment. Our analysis showed that fatty acid metabolic process MT correlated with significantly increased natural killer T (NKT) cells and significantly decreased CD8+T cells. At the same time, GSEA analysis revealed that fatty acid metabolic process MT was associated with significantly increased immune activation pathways and an enriched fraction of cytokine secretion compared with WT.
**INTRODUCTION**

Even if a glioblastoma (GBM), a primary brain tumor with the highest degree of intracranial malignancy, can be safely and fully resected, the median survival time for patients is still only 14 months, and there is recurrence rate of 90% (1, 2). That being said, the main treatment of early GBM is still surgical resection. Radiotherapy and chemotherapy are also important methods for treating GBM, and some clinical studies have shown that these treatments can play an active role in improving the survival time of patients. However, these methods are not without their drawbacks (3, 4). Normal brain tissue damage is caused by the low selectivity of radiotherapy and chemotherapy, while the use of chemotherapy drugs can cause systemic immunosuppression. In addition, because of the complexity and dynamic changes of tumor microenvironments, chemotherapy drugs are prone to drug resistance (4–7).

In recent years, there has been a succession of breakthroughs in the field of tumor immunotherapy. During this time, the Food and Drug Administration (FDA) has approved a number of new immunotherapy drugs for various types of tumor therapies, such as immune-checkpoint inhibitors (ICIs) and chimeric antigen receptor (CAR) T-cells (8). At present, there are many studies exploring the application effect of GBM immunotherapy. These studies have shown that in the GBM mouse model, the therapeutic inhibition of IDO, cytotoxic T lymphocyte antigen-4 (CTLA-4), or programmed death-ligand 1 (PD-L1) can significantly reduce the proportion of activated immune cells, and improve the immune microenvironment.

**Conclusions:** We found that fatty acid metabolic process MT may be used as an independent predictor of the efficacy of ICI treatment in GBM patients. Use of the fatty acid metabolic process MT will result in higher immunogenicity rates, a significant increase in the proportion of activated immune cells, and improvement of the immune microenvironment.

**Keywords:** glioblastoma, immune-checkpoint inhibitors, immunotherapy, prognosis, fatty acid metabolic process

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**METHODS**

**GBM Queue**

Samstein et al. published a GBM queue (ICI-GBM) that accepts ICIs, including mutation data analyzed via targeted sequencing and survival data (18). Due to the limited number of GBM queues for immune checkpoint treatment, we also collected TCGA-GBM queues (19) from the GDC database and a Jonsson-GBM queue (20) from cBioPortal. The TGCA-GBM queues included expression data, mutation data, and survival data (20). The Jonsson-GBM queue included mutation data and survival data. See Table S1 for details on the clinical features of patients in the ICI-GBM cohort.

**Mutation State of the Fatty Acid Metabolic Process Pathway**

In order to explore the relationship between the mutation status of the fatty acid metabolic process pathway and the clinical prognosis of GBM patients after ICI treatment, we downloaded the gene set related to the fatty acid metabolic process from the Molecular Signatures Database (MsigDB) (https://www.gsea-msigdb.org/gsea/msigdb/genesets.jsp) (21) database in the format of a GMT file. First, we screened the mutation data in the ICI-GBM, TCGA-GBMs, and Jonsson-GBM queues based on the definition of non-synonymous mutation types in the maftools R package (22). After this initial screening, only non-synonymous mutation data was retained for subsequent analysis. From here, we calculated the mutation rate of each patient’s fatty acid metabolic process and defined the mutation state of each patient’s pathway according to whether any mutation occurred.
If the number of mutations in this pathway was zero, it was considered a WT. If the number of mutations in this pathway was greater than 0, it was considered an MT.

**Immune Microenvironment Analysis and Drug Sensitivity Analysis**

In order to explore the difference in the immune cell scores between the WT and MT groups, we used CIBERSORT and IOBR to evaluate the immune cell scores of each patient (23). In addition, we used ssGSEA and GSEA algorithms to evaluate the activation degree of patients in each pathway (24, 25). The emapplot function was used in our enrichment analysis to analyze the expression data of each patient (26). From this, we obtained the estimated IC50 value of each patient for 138 drugs from the Genomics of Drug Sensitivity in Cancer (GDSC) database (https://www.cancerrxgene.org/) (27).

**Statistical Analysis**

To explore the relationship between the fatty acid metabolic process and the immunotherapy prognosis of GBM patients, we used Kaplan Meier (KM) survival analysis, univariate and multivariate Cox regression analysis, and log-rank P to show the degree of statistical difference. In the difference analysis of continuous variables, the Wilcoxon-Mann-Whitney test was used to compare the amount of tumor mutation burdens (TMB), DNA damage repair (DDR) pathway mutations, and the immune cell scores. To estimate the difference of IC50 between the WT and MT groups, Fisher's exact test was used to count the difference in the mutation frequency between each group. The difference was analyzed using the Limma Package (28). GSEA and enrichment analysis were performed using Clusterprofiler (26). A heat map was drawn using the Complexheatmap package (29). Box drawings were created using ggplot2 (30). The judging range of differential genes in the analysis was |log2FC|>1 and P < 0.05. The analysis flow of this study is shown in Figure 1.

**RESULTS**

**Prognostic Role of the Mutation State of the Fatty Acid Metabolic Process in GBM Treatment**

In order to explore the relationship between the fatty acid metabolic process and the immunotherapy prognosis of GBM patients, univariate and multivariate Cox regression analysis were applied to the ICI-GBM cohort. Univariate results showed that fatty acid metabolic process MT was related to significantly prolonged OS time in GBM patients (Figure 2A; P < 0.05; HR = 0.39). Through our analysis, we found that fatty acid metabolic process MT can be used as an independent predictor of ICIs in GBM patients (Figure 2B; P < 0.05; HR = 0.34). In ICI-GBM queue, survival analysis showed that fatty acid metabolic process MT had significantly longer OS time than fatty acid metabolic process WT [Figure 2C; P = 0.033; Hazard Ratio (HR) = 0.4; 95% CI: 0.22 - 0.75]. We also found that the mutation status of the fatty acid metabolic process can predict other immune queues. For example, in ICI-Bladder cancer, survival analysis showed that fatty acid metabolic process MT resulted in significantly longer OS time than fatty acid metabolic process WT (Figure 2D; P = 0.019; HR = 0.5; 95% CI: 0.31 – 0.81). In TCGA-GBM, the mutation state of fatty acid metabolic process has nothing to do with the prognosis state of GBM (Figure 2E; P > 0.05). Similarly, in Jonsson-GBM, the OS time and progression-free survival (PFS) time of patients had nothing to do with the mutation state of the fatty acid metabolic process (Figures 2F, G).

**The Mutation State of the Fatty Acid Metabolic Process and Immune Cells**

Firstly, we analyzed the differences in expression data between fatty acid metabolic process MT and WT. In Figure 3A, the genes that were significantly up-regulated and significantly down-regulated in the WT and MT groups are displayed in red and blue, respectively (P < 0.05; |log2FC| > 1). Based on immune-related genes and the ssGSEA algorithm, we found that the fraction of NK cells in the MT group was significantly higher.
than that in WT group. On the other hand, we also found that the level of depletion among the CD8+T cells, the scores of co-inhibition antigen-presenting cells (APC), and the number of co-inhibition T cells in the MT group were significantly lower than those in WT group (Figure 3B; P < 0.05). In addition, using the CIBEROSRT algorithm to evaluate the expression data of patients, we found that the proportion of memory B cells in the MT group was significantly higher than that in the WT group (Figure 3C; P < 0.05). On the other hand, the relative ratio of resting memory CD4+ T cells in the MT group was significantly higher than that in WT group (Figure 3D; P < 0.05).

The Mutation State and Immunogenicity of the Fatty Acid Metabolic Process

We began by analyzing the relationship between TMB and the mutation state of the fatty acid metabolic process. We found that the level of TMB in the MT group was significantly higher than that in the WT group (Figure 4A; P < 0.05). In addition, compared with the
In this study, univariate and multivariate Cox regression analyses showed that fatty acid metabolic process MT may be an independent prognostic factor for GBM patients receiving ICI treatment. In addition, Kaplan Meier survival analysis showed that the MT group had significantly longer OS time than the WT group.
The results of immune microenvironment analysis showed that the MT group had an inflammatory immune microenvironment that manifested as significantly up-regulated NKT cells and significantly down-regulated exhausted CD8+ T cells. The results showed that the activation of immune cells, as well as the secretion and production pathways of inflammatory molecules, were significantly activated in the MT group. At the same time, the MT group displayed a significant increase in immunogenicity compared to the WT group.

The higher immunogenicity of patients with fatty acid metabolic process MT is potentially the mechanism behind patients’ better prognoses post-ICI treatment. Patients with fatty acid metabolic process MT not only had higher TMB, but also a higher number of gene mutations in their DDR pathway, which improved their immunogenicity. Due to gene fusion, deletion mutation, and point mutation of tumor genes, new abnormal proteins, which are new antigens encoded by the mutant genes of tumor cells, are produced (31). These new tumor antigens can not only enhance the immune response of T cells, but also increase the content of new antigen-specific T cells, thus improving the immune system’s ability to identify tumors (32). Studies have shown that the mutation of the DDR pathway will result in the production of higher TMB and neoantigen loads, thus significantly improving the effectiveness of ICI treatment (33–35).

An inflammatory immune microenvironment also results in greater effectiveness for ICIs treatment in patients with fatty acid metabolic process. The MT group showed a significant increase in activated immune cells such as NK cells and a significant decreased in depleted CD8+ T cells. In these groups, we found that NKT cells...
killed tumor cells in three ways and thus played an important anti-tumor role (1): in the Fas/FasL, perforin, and granzyme B pathways, they played a cytotoxic role and directly kill tumor cells (36)(2); they regulate the recruitment and function of other immune cells by secreting cytokines, thus indirectly playing an anti-tumor role (3); NKT cells changed the immunosuppression level in the immune microenvironment, which resulted in anti-tumor activities (38). In addition, when CD8+T cells bind to antigens from dendritic cells, cytotoxic CD8+T cells can be produced. Under the induction of chemokines secreted by the dendritic cells CXCL9 and CXCL10, activated cytotoxic CD8+T cells were able to migrate to the inflammatory environment through CXCR3 expression (39). In the immune microenvironment, IL-1 helped to establish a pro-inflammatory environment by inducing pro-inflammatory cytokines and chemokines (40). IFN-γ played a central role in immunity by inducing MHCI molecules, Fas/FasL, and inducing immune proteasome-related factors (41). In the IFN-γ environment, the main cytokines secreted by activated CD8+T cells significantly increased the expression of PD-L1 (42).

This study has some limitations. Firstly, the number and sample size of clinical studies of glioma patients treated with immune checkpoint inhibitors in the database are currently small, so we hope to recruit more GBM patients who have received ICI therapy in the future to further verify the conclusions of this study. Second, few animal or cell experiments have been done on the subject. Third, the ICI-GBM cohort contained only the mutation data and clinical prognosis data of targeted sequencing, so there may have been

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**FIGURE 5**

(A) The gene sets enrichment analysis showed that the enrichment scores of the immune cells were significantly up-regulated in the fatty acid metabolic process of the MT group. (B) The gene sets enrichment analysis showed that the enrichment scores of cytokine production were significantly up-regulated in the fatty acid metabolic process of the MT group. (C) The emapplot representing the connection between the immune-related signaling pathways. MT, mutant-type.
potential bias when calculating the number of gene mutations in the fatty acid metabolic process pathway.

CONCLUSIONS

In this study, we found that fatty acid metabolic process MT may be used as an independent predictor of the efficacy of ICI treatment in GBM patients, which will allow doctors to better screen out the dominant population receiving immunotherapy in the future. Fatty acid metabolic process MT is related to higher immunogenicity, a significant increase in the quantity of activated immune cells, and improvement to the immune microenvironment.

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.

ETHICS STATEMENT

Ethical review and approval was not required for the study on human participants in accordance with the local legislation and institutional requirements. Written informed consent for

FIGURE 6 | (A) A comparison of the estimated IC50 values of methotrexate in the fatty acid metabolic processes of the MT and WT groups. (B) The results of the DNA binding signaling GSEA. (C) A comparison of the estimated IC50 values of pictilisib in the fatty acid metabolic processes of the MT and WT groups. (D) The GSEA results of PI3K/AKT signaling. MT, mutant-type; WT, wild-type; IC50, half maximal inhibitory concentration; GSEA, Gene Set Enrichment analysis. *P < 0.05; **P < 0.01.
participation was not required for this study in accordance with the national legislation and the institutional requirements.

AUTHOR CONTRIBUTIONS

Conceptualization, BL and ZQ. Formal analysis, RL, WL, QH, LW. Visualization, RL, QL. Writing–original draft, RL, WL, QH, LW, FZ, BL and ZQ. Writing–review & editing, RL, WL, QH, LW, FZ, BL and ZQ. All authors read and approved the final manuscript.

FUNDING

This work was supported by the Science and Technology Project of Education Department of Jiangxi Province (180805), the Science and Technology Project of Health and Family Planning Commission of Jiangxi Province (20195363), the CSCO-Haosen Oncology Research Fund (Y-HS2019/2-015), and the Science and Technology Project of Ganzhou City (GZ2021SF002).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.819515/full#supplementary-material

Supplementary Table 1 | The clinical characteristics of the ICI-GBM cohort.

Supplementary Figure 1 | The co-exclusivity and co-occurrence of top 30 mutated driver genes in the fatty acid metabolic processes of the WT (A) and MT (B) groups. MT, mutant-type; WT, wild-type.

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