

GPR27 expression correlates with prognosis and tumor progression in gliomas

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Backgrounds: Glioma is a highly aggressive type of brain tumor, and its prognosis is still poor despite recent progress in treatment strategies. G protein-coupled receptor 27 (GPR27) is a member of the G protein-coupled receptor family and has been reported to be involved in various cellular processes, including tumor progression. Nevertheless, the clinical potential and tumor-related role of GPR27 in glioma remain unknown. Here we aimed to explore the function and role of GPR27 in gliomas.

Methods: In the current study, we evaluated the expression and clinical significance of GPR27 in gliomas using data from GTEx and TCGA datasets. We also conducted cellular experiments to evaluate the functional role of GPR27 in glioma cell growth.

Results: We found that glioma tissues had significantly lower GPR27 expression levels than normal brain specimens, and its expression level was closely associated with disease status. Of note, GPR27 was negatively correlated with WHO grade, with grade IV samples showing the lowest GPR27 levels, while grade II samples showed the highest levels. Patients with IDH mutation or 1p/19q co-deletion exhibited higher GPR27 levels. In addition, lower GPR27 levels were correlated with higher death possibilities. In cellular experiments, we confirmed that GPR27 inhibited glioma cell growth.

Conclusions: Our results indicate that GPR27 may function as a potential prognostic biomarker and therapeutic target in gliomas. Further studies are needed to illustrate the signaling mechanism and clinical implications of GPR27 in gliomas.

1 **GPR27 expression correlates with prognosis and tumor progression in gliomas**

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17 **Data availability:** Original data was provided and uploaded.

18 **Ethics**

19 The Ethics Committee of Suining Central Hospital requires no ethic approval and grants an
20 exemption from informed consent for this public-database study.

21

22 **Abstract**

23 **Backgrounds:** Glioma is a highly aggressive type of brain tumor, and its prognosis is still poor
24 despite recent progress in treatment strategies. G protein-coupled receptor 27 (GPR27) is a
25 member of the G protein-coupled receptor family and has been reported to be involved in various
26 cellular processes, including tumor progression. Nevertheless, the clinical potential and tumor-
27 related role of GPR27 in glioma remain unknown. Here we aimed to explore the function and role
28 of GPR27 in gliomas.

29 **Methods:** In the current study, we evaluated the expression and clinical significance of GPR27
30 in gliomas using data from GTEx and TCGA datasets. We also conducted cellular experiments to
31 evaluate the functional role of GPR27 in glioma cell growth.

32 **Results:** We found that glioma tissues had significantly lower GPR27 expression levels than
33 normal brain specimens, and its expression level was closely associated with disease status. Of
34 note, GPR27 was negatively correlated with WHO grade, with grade IV samples showing the
35 lowest GPR27 levels, while grade II samples showed the highest levels. Patients with IDH
36 mutation or 1p/19q co-deletion exhibited higher GPR27 levels. In addition, lower GPR27 levels
37 were correlated with higher death possibilities. In cellular experiments, we confirmed that GPR27
38 inhibited glioma cell growth.

39 **Conclusions:** Our results indicate that GPR27 may function as a potential prognostic biomarker
40 and therapeutic target in gliomas. Further studies are needed to illustrate the signaling
41 mechanism and clinical implications of GPR27 in gliomas.

42

43 **Keywords**

44 Biomarkers, Cell proliferation, Glioblastoma, Neuroscience, Molecular biology, Survival analysis.

45

46 Introduction

47 Gliomas and glioblastomas are types of primary brain tumors that originate from glial cell, which
48 are the supportive cells of the central nervous system [1]. Glioma is the most common form of
49 brain malignancies, accounting for about 80% of brain tumors, and they can vary in grade from
50 slow-growing low-grade tumor to aggressive high-grade tumor. Among gliomas, glioblastoma,
51 also known as glioblastoma multiforme (GBM), represents the most aggressive subtype,
52 representing the highest grade of glioma [2]. Glioblastoma is characterized by its rapid growth,
53 infiltrative nature, and resistance to treatment, making it one of the most challenging cancers to
54 manage. Understanding the basic features and biology of gliomas and glioblastomas is crucial
55 for advancing diagnosis, prognosis, and treatment strategies for these complex brain tumors [3].

56 *GPR27*, a member of the G protein-coupled receptor family, is a protein-coding gene located on
57 human chromosome 3q25.1 [4]. It encodes a 7-transmembrane receptor that expresses in
58 various tissues and is involved in multiple cellular functions including neurotransmission, immune
59 activation, and cellular growth [5, 6]. Recent research has implied that GPR27 might play critical
60 roles during cancer progression, particularly in hepatocellular carcinoma (HCC) [7] and breast
61 cancer [8]. Abnormal expression of GPR27 has been observed in these cancers, and it has been
62 recognized to contribute to tumor growth and angiogenesis. For example, Wang et al. discussed
63 function of GPR27 in HCC progression and its potential mechanism of action through MAPK-
64 ERK signaling pathway [7]. The study highlights the importance of GPR27 in HCC, a type of liver
65 cancer, and proposes that GPR27 may promote tumor progression by activating the MAPK/ERK
66 pathway. The paper provides insights into the potential molecular mechanisms underlying HCC
67 development and indicates that GPR27 could be a valuable therapeutic target in HCC treatment.
68 However, further study will be needed to fully understand the exact mechanisms by which
69 GPR27 affects cancer development and to explore its potential as a treatment target for cancers.
70 This study sheds light on expression and clinical relevance of GPR27 in gliomas and provides
71 evidence for its oncogenic role in glioma growth. The findings presented here have the potential

72 to enhance our understanding of underlying mechanism of glioma progression and could
73 contribute to the development of novel therapeutic strategies for this devastating disease.

74

75 **Methods**

76 **In silico data collection and analysis**

77 RNA expression information in the Fragments Per Kilobase per Million format (FPKM) as well as
78 related clinic-pathological data of glioma samples (n=689) and normal brain samples (n=1157)
79 were downloaded from TCGA (<https://portal.gdc.cancer.gov/projects/TCGA-GBM>) and GTEx
80 (<https://www.gtexportal.org/home/>). The clinic-pathological information of glioma patients was
81 also extracted. The difference of various clinicopathological parameters were compared between
82 the high-GPR27 and low-GPR27 expression groups. Regression analyses were used to assess
83 the relationships between GPR27 expression and clinicopathological variables of glioma cases.

84 Survival information of glioma cases in TCGA-glioma was analyzed. Kaplan-Meier survival as
85 well as multivariate Cox regression analyses was conducted to identify the patients' prognoses
86 according to GPR27 level as well as other clinic-pathological characteristics.

87 **Establishment of nomogram and calibration curves**

88 We employed the "RMS" package in R to construct the nomogram for predicting possibility of
89 individual survival. The calibration of the nomogram was evaluated through calibration curves.

90 **Immune infiltration analyses**

91 We evaluated the association between GPR27 expression and immune cell infiltration in gliomas
92 using ssGSEA algorithm available in the "GSVA" R package [9]. This algorithm helped us
93 assess infiltration status of 24 kinds of immune cell types [10]. We performed Spearman
94 correlation analysis to clarify correlation between GPR27 expression and immune cell infiltration
95 status.

96 **Cell culture and transfection**

97 U87 and U251 human glioblastoma cells will be obtained from ATCC and cultured in Dulbecco's
98 Modified Eagle's Medium (DMEM) supplemented with 10% FBS and 1% penicillin-streptomycin
99 at 37°C with 5% CO₂. Culturing medium was replaced every two days, and cells were sub-
100 seeded once they reach 70-80% confluence. Transient transfection of siRNAs targeting GPR27
101 or scramble control siRNA was achieved with Lipofectamine 3000 based on manufacturer's
102 instructions. Afterwards, cells were incubated to allow for transfection. After the incubation period,
103 the transfection medium will be replaced with fresh complete growth medium to allow for further
104 cell culture or further experimental tests.

105 **Cell proliferation assay**

106 Transfected cells were seeded at 5000 cells/well. Then CCK-8 experiment was done according
107 to the manufacturer's instruction. Briefly, the medium was replaced with CCK-8 reagent, and the
108 cells were incubated for 2 hours. The absorbance of the formazan dye, which is proportional to
109 the number of viable cells, was measured spectrophotometrically at 450 nm. The percentage of
110 cell viability or proliferation was calculated by normalizing to the control group. Results were
111 expressed as mean \pm SD to determine statistical significance.

112 **Statistics**

113 All statistical analyses and plots were performed using R (version 4.1.3). Statistical significance
114 was set at $P < 0.05$. * indicates $P < 0.05$, ** indicates $P < 0.01$, *** indicates $P < 0.001$.

115 **Ethics**

116 The Ethics Committee of Suining Central Hospital requires no ethic approval and grants an
117 exemption from informed consent for this public-database study.

118

119 **Results**

120 **Aberrant GRP27 expression in gliomas**

121 As shown in Figure 1A, RNA expression data from GTEx and TCGA datasets suggested that
122 glioma tissues possessed a significant lower GPR27 level than normal brain tissues ($P < 0.001$).
123 Meanwhile, GPR27 expression was closely correlated with disease status. For example, GPR27
124 was negatively correlated with WHO grade, on that grade IV samples showed the lowest GPR27
125 level while grade II samples showed the highest GPR27 level (Figure 1B, $P < 0.001$). In contrast,
126 patients with IDH mutation (Figure 1C) or 1p/19q co-deletion (Figure 1D) exhibited higher
127 GPR27 levels ($P < 0.001$). Moreover, *in silico* analysis revealed that lower-GPR27 level was
128 correlated with higher death rates (Figure 1E, $P < 0.001$) although its prognostic significance
129 remains further investigation.

130 **Correlations between GPR27 expression and clinic-pathological characteristics of** 131 **gliomas**

132 By dividing TCGA cohorts into high-GPR27 group ($n=350$) and low-GPR27 group ($n=349$), we
133 further analyzed its correlation with clinic-pathological variables (Table 1). Accordingly, lower
134 GPR27 expression was observed in elder patients, higher-grade gliomas, as well as in the
135 specimens with wild-type IDH status or non-codeletion of 1p/19q (all $P < 0.001$). Considering all
136 the above-mentioned variables had been reported to be correlated with patients' prognosis, we
137 were engaged to further explore whether GPR27 can affect the overall survival of glioma patients.
138

139 **GPR27 is an independent survival predictor of gliomas**

140 Kaplan-Meier survival analysis showed that patients with lower GPR27 level had significantly
141 worse prognosis (Figure 2A, $P < 0.001$). Briefly, the median overall survival time of low-GPR27
142 group was 26.3 months (95% CI 23.6-34.1 months), while was up to 99.6 months (95% CI 68.4-
143 135.6 months) of high-GPR27 group. In addition to GPR27, several other variables were
144 identified with prognostic significances in univariate analyses. For example, comparing with
145 WHO grade II patients, WHO grade III patients showed a death hazard ratio as 2.967 (95% CI
146 1.986 - 4.433, $P < 0.001$) and WHO grade IV patients showed a death hazard ratio as 18.6 (95%

147 CI 12.448 - 27.794, $P < 0.001$). Consistent with our previous data, mutated IDH status indicated a
148 better survival with a death hazard ratio as 0.116 (95% CI 0.089 - 0.151, $P < 0.001$) comparing to
149 those with wild-type IDH status. Similarly, co-deletion of 1p/19q was a favorable prognostic factor
150 with a death hazard ratio as 0.225 (95% CI 0.147 - 0.346, $P < 0.001$).

151 To figure out the independent prognostic factors, we further subjected the variables into a Cox
152 multivariate regression model for survival analysis (Table 2). As a result, elder age and higher
153 WHO grade were confirmed as independent unfavorable prognosis factors. In contrast, mutated
154 IDH status was identified as an independent favorable prognostic factor (hazard ratio 0.265, 95%
155 CI 0.176 - 0.399, $P < 0.001$). Importantly, our data, for the first time, showed that higher GPR27
156 expression was an independent benefit biomarker for glioma prognosis (hazard ratio 0.679, 95%
157 CI 0.486 - 0.947, $P = 0.023$).

158 Based on the multivariate survival analysis, we also established a nomogram to help predict
159 overall survival of glioma patients (Figure 2B); the variables in the nomogram included patients'
160 age, gender, WHO grade, IDH status, 1p/19q co-deletion, and GPR27 expression level.

161 **GPR27 inhibits proliferation of glioma cells**

162 We next conducted western blot experiments to confirm the efficiency of knocking down GPR27
163 in U87 and U251 cell lines. The data obtained from the western blot experiments demonstrated a
164 significant reduction in GPR27 expression levels, which indicated that the knockdown was
165 successful (Figure 3A). To further investigate the role of GPR27 in cell proliferation, we
166 conducted CCK-8 experiments. The results showed that silencing GPR27 significantly inhibited
167 the proliferation (Figure 3B- 3C) capacities of both U87 and U251 cell lines. These findings
168 suggested that GPR27 plays a tumor-promoting role in glioma cells. The inhibition of GPR27
169 expression may be a promising therapeutic approach for treating glioma. Overall, the results of
170 this study provide important insights into the molecular mechanisms underlying glioma
171 development and progression, and may have significant implications for the development of
172 novel therapeutic strategies for this devastating disease.

173 **GPR27 is correlated with the immune cell infiltration in gliomas**

174 The results of GSEA showed that the GPR27 has a negative association with macrophages (R=-
175 0.551, $P < 0.001$), neutrophils (R=-0.473, $P < 0.001$), aDC (R=-0.405, $P < 0.001$), eosinophils (R=-
176 0.397, $P < 0.001$), iDC cells (R=-0.324, $P < 0.001$), cytotoxic cells (R=-0.286, $P < 0.001$). etc. These
177 findings indicate that GPR27 may play a crucial role in modulating the immune response in the
178 tumor microenvironment. Oppositely, GPR27 shows positive associations with pDC cells
179 (R=0.325, $P < 0.001$), TFH cells (R=0.281, $P = 0.003$), NK CD56bright cells (R=0.251, $P < 0.001$),
180 Tcm cells (R=0.214, $P < 0.001$), TReg cells (R=0.190, $P < 0.001$), etc (Figure 4A-C). These results
181 suggest that GPR27 may play a significant role in promoting the differentiation and activation of
182 these immune cells.

183

184 **Discussions**

185 The results presented in this study suggest that GPR27 expression is significantly reduced in
186 glioma tissues when compared to normal brain tissues and that GPR27 expression levels are
187 closely correlated with the disease status of glioma patients. Specifically, GPR27 expression was
188 negatively correlated with WHO grade, and patients with IDH mutation or 1p/19q co-deletion
189 exhibited higher GPR27 levels. Additionally, in silico analysis showed that lower GPR27
190 expression was correlated with higher death rates in glioma patients. In line with these findings,
191 the authors performed further analyses and showed that low GPR27 expression was associated
192 with elder patients, higher-grade gliomas, and wild-type IDH status or non-codeletion of 1p/19q.
193 Importantly, Kaplan-Meier survival analysis indicated that lower GPR27 expression was a
194 significant independent predictor of poor overall survival in glioma patients. Taken together,
195 clinical data analyses suggest that GPR27 may serve as a useful prognostic biomarker for
196 glioma and may have implications for the development of novel therapeutic strategies.

197 Consistently, cellular experiments suggest that GPR27 plays a crucial role in the development
198 and progression of glioma, as silencing of GPR27 expression significantly inhibited the

199 proliferation capacities of U87 and U251 cell lines. These findings are consistent with previous
200 reports that have demonstrated that GPR27 is overexpressed in several types of cancer and
201 promotes tumor growth and metastasis. For example, Wang et al. confirmed that GPR27
202 expression was upregulated in HCC tissues and cell lines. They then used cellular experiments
203 to show that knockdown of GPR27 inhibited HCC cell proliferation, migration, and invasion.
204 Further analysis revealed that GPR27 acted through the MAPK/ERK pathway to promote HCC
205 progression [7]. Besides the classical downstream G-protein signaling, GPR27 can also
206 activate beta-arrestin-biased signaling pathways [6]. Meanwhile, beta-arrestins had been well-
207 acknowledged to be involved in malignancies including glioblastoma [11, 12]. Therefore, its high
208 likely that GPR27 may promotes glioma progression via beta-arrestin-downstream signaling
209 pathways. The successful knockdown of GPR27 in this study confirms the specificity of the effect
210 observed on cell proliferation. Furthermore, the identification of GPR27 as a potential therapeutic
211 target for glioma treatment is of significant clinical relevance, as there is a dire need for novel
212 therapeutic strategies for this devastating disease. Further studies will be necessary to further
213 explore its functional mechanisms.

214 The present study provides important insights underlying glioma development and progression.
215 The observed negative correlation between GPR27 and various immune cell types suggests that
216 GPR27 may play a role in immune evasion by glioma cells, and further investigations into the
217 mechanism underlying this association could help to develop effective immunotherapeutic
218 strategies [13, 14]. Moreover, the identification of GPR27 as a potential therapeutic target could
219 pave the way for the development of new drugs that selectively inhibit GPR27 expression or
220 function, which could offer a promising avenue for glioma treatment. However, more research is
221 needed to fully understand the mechanisms underlying the role of GPR27 in glioma development
222 and progression, as well as the potential therapeutic implications of targeting GPR27.

223 Furthermore, our data identified significant correlations between GPR27 and different immune
224 cells. The implications of these findings could be far-reaching in the field of cancer
225 immunotherapy. Given the crucial role that the tumor microenvironment plays in cancer

226 progression [15], GPR27 could serve as a potential target for the development of novel cancer
227 immunotherapies. By inhibiting the expression of GPR27, it may be possible to modulate the
228 immune response in the tumor microenvironment to promote an anti-tumor immune response.
229 Alternatively, targeting GPR27 could be used to promote the differentiation and activation of
230 immune cells that are positively associated with GPR27, such as pDC cells and TReg cells.
231 Overall, these findings provide important insights into the role of GPR27 in the modulation of the
232 immune response in the tumor microenvironment and could have significant implications for the
233 development of novel cancer immunotherapies.

234 The study has several limitations, such as a restricted sample size, which may not be
235 representative of the entire population. Moreover, the study's outcomes may not be generalized
236 to other regions due to its geographic confinement. The self-reported data may also be
237 influenced by potential biases. Furthermore, the study's cross-sectional design prevents
238 establishing a cause-and-effect relationship between variables.

239

240 **Conclusions**

241 Taken together, our data highlights the potential clinical significance of GPR27 in gliomas. The
242 findings indicate that GPR27 expression is significantly lower in glioma tissues than in normal
243 brain tissues and is negatively correlated with WHO grade. Additionally, lower GPR27
244 expression is associated with elder patients, higher-grade gliomas, and specimens with wild-type
245 IDH status or non-codeletion of 1p/19q, and predicts worse prognosis. These results suggest
246 that GPR27 may be a potential prognostic biomarker for glioma patients. However, further
247 research is needed to fully understand the role of GPR27 in gliomas and to explore its potential
248 as a therapeutic target.

249

250 **Acknowledgement**

251 None.

252

254 **Figure legends**

255 **Figure 1. The correlation between GPR27 expression and disease status in glioma.**

256 A) RNA level data from GTEx and TCGA datasets revealed that glioma tissues had significantly
257 lower GPR27 expression levels than normal brain tissues ($P < 0.001$).

258 B) GPR27 expression was negatively correlated with WHO grade, with grade IV samples
259 exhibiting the lowest GPR27 levels and grade II samples showing the highest levels ($P < 0.001$).

260 C) Patients with IDH mutation showed higher GPR27 levels ($P < 0.001$).

261 D) Similarly, patients with 1p/19q co-deletion exhibited higher GPR27 levels ($P < 0.001$).

262 E) In silico analysis indicated that lower GPR27 expression levels were correlated with higher
263 death rates ($P < 0.001$).

264

265 **Figure 2. Prognostic significance of GPR27 on glioma patients.**

266 A) Kaplan-Meier survival analysis of glioma patients based on GPR27 expression level. Patients
267 with lower GPR27 level had significantly worse prognosis compared to those with higher GPR27
268 level ($P < 0.001$).

269 B) A nomogram established based on the multivariate survival analysis. The variables in the
270 nomogram included patients' age, gender, WHO grade, IDH status, 1p/19q co-deletion, and
271 GPR27 expression level.

272

273 **Figure 3. GPR27-knockdown inhibit glioma progression.**

274 A) Western blot analysis confirmed the successful knockdown of GPR27 expression in U87 and
275 U251 cell lines.

276 B) CCK-8 assay results indicated that silencing GPR27 significantly inhibited the proliferation of
277 U87 cells compared to the control group.

278 C) CCK-8 assay results indicated that silencing GPR27 significantly inhibited the proliferation of
279 U251 cells compared to the control group.

280

281 **Figure 4. The correlation between immune cell infiltration and GPR27 expression in**
282 **glioma.**

283 A) Spearman analysis result showed the correlation between the infiltration of 24 types of
284 immune cells and GPR27 expression in glioma tissues.

285 B) and C) showed representative infiltration level of pDC and macrophages in cells with different
286 GPR27 level.

287 DC, dendritic cells; pDC, plasmacytoid DC; aDC, activated DC; iDCs, immature DCs; NK, natural
288 killer cells; Tgd, T gamma delta; TReg, regulatory T cells; Tem, T effector memory; Tcm, T
289 central memory; Th1 cells, type 1 Th cells; Th2 cells, type 2 Th cells; Th17 cells, type 17 Th cells;
290 TFH, T follicular helper.

291

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326

Table 1 (on next page)

Table 1. Basic information of enrolled glioma patients.

1 Table 1. Basic information of enrolled glioma patients.
2

3	Characteristics	Low GPR27	High GPR27	P value
4	Total cases, n	349	350	
5	Age, n (%)			< 0.001***
6	≤ 60 years old	245 (35.1%)	311 (44.5%)	
	> 60 years old	104 (14.9%)	39 (5.6%)	
	Gender, n (%)			0.134
	Female	139 (19.9%)	159 (22.7%)	
	Male	210 (30%)	191 (27.3%)	
	WHO grade, n (%)			< 0.001***
	G2	67 (10.5%)	157 (24.6%)	
	G3	119 (18.7%)	126 (19.8%)	
	G4	141 (22.1%)	27 (4.2%)	
	IDH status, n (%)			< 0.001***
	WT	189 (27.4%)	57 (8.3%)	
	Mutation	152 (22.1%)	291 (42.2%)	
	1p/19q codeletion, n (%)			< 0.001***
	Non-codel	328 (47.4%)	192 (27.7%)	
	Codel	14 (2%)	158 (22.8%)	
	OS event, n (%)			< 0.001***
	Alive	162 (23.2%)	265 (37.9%)	
	Dead	187 (26.8%)	85 (12.2%)	

Table 2 (on next page)

Table 2. Univariate and multivariate Cox regression analyses of the overall survival of glioma patients.

1 Table 2. Univariate and multivariate Cox regression analyses of the overall survival of glioma patients.
2

Characteristics	Total (N)	Univariate analysis		Multivariate analysis	
		Hazard ratio (95% CI)	P value	Hazard ratio (95% CI)	P value
Age	698				
≤ 60 years old	555	Reference		Reference	
> 60 years old	143	4.696 (3.620 - 6.093)	< 0.001***	1.429 (1.046 - 1.952)	0.025*
Gender	698				
Female	297	Reference		Reference	
Male	401	1.250 (0.979 - 1.595)	0.073	1.185 (0.903 - 1.553)	0.220
WHO grade	636				
G2	223	Reference		Reference	
G3	245	2.967 (1.986 - 4.433)	< 0.001***	1.873 (1.223 - 2.868)	0.004**
G4	168	18.600 (12.448 - 27.794)	< 0.001***	4.253 (2.525 - 7.162)	< 0.001***
IDH status	688				
WT	246	Reference		Reference	
Mutation	442	0.116 (0.089 - 0.151)	< 0.001***	0.265 (0.176 - 0.399)	< 0.001***
1p/19q codeletion	691				
Non-codel	520	Reference		Reference	
Codel	171	0.225 (0.147 - 0.346)	< 0.001***	0.843 (0.494 - 1.440)	0.533
GPR27	698				
Low	349	Reference		Reference	
High	349	0.326 (0.251 - 0.423)	< 0.001***	0.679 (0.486 - 0.947)	0.023*

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4
5
6
7
8

Figure 1

Figure 1. The correlation between GPR27 expression and disease status in glioma.

A) RNAseq data from GTEx and TCGA datasets revealed that glioma tissues had significantly lower GPR27 expression levels than normal brain tissues ($P < 0.001$). B) GPR27 expression was negatively correlated with WHO grade, with grade IV samples exhibiting the lowest GPR27 levels and grade II samples showing the highest levels ($P < 0.001$). C) Patients with IDH mutation showed higher GPR27 levels ($P < 0.001$). D) Similarly, patients with 1p/19q co-deletion exhibited higher GPR27 levels ($P < 0.001$). E) In silico analysis indicated that lower GPR27 expression levels were correlated with higher death rates ($P < 0.001$).

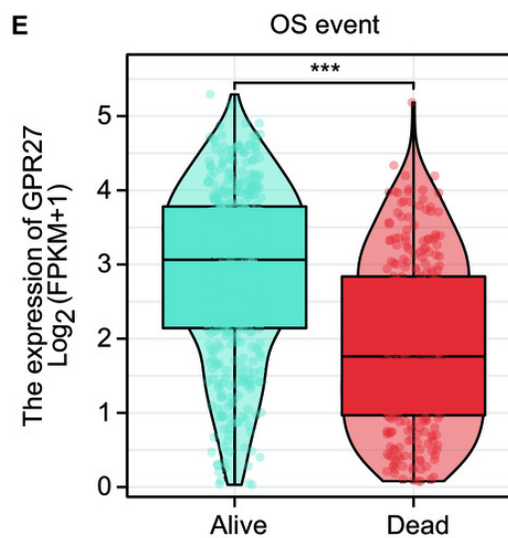
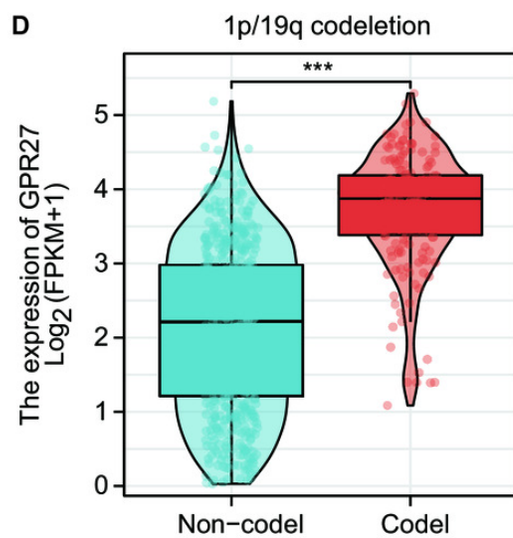
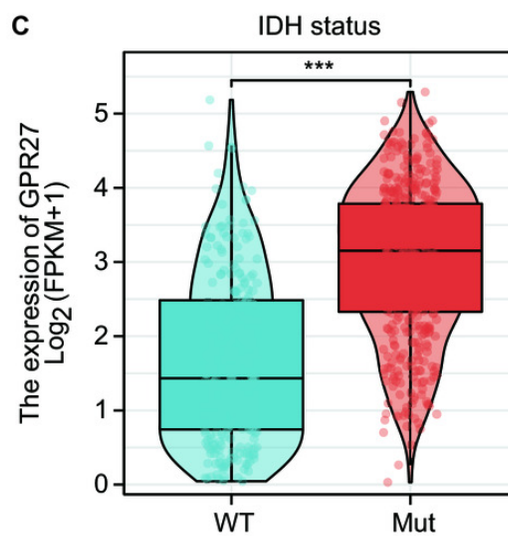
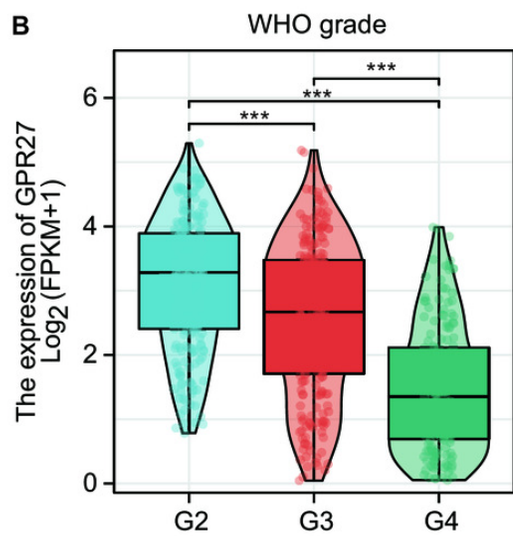
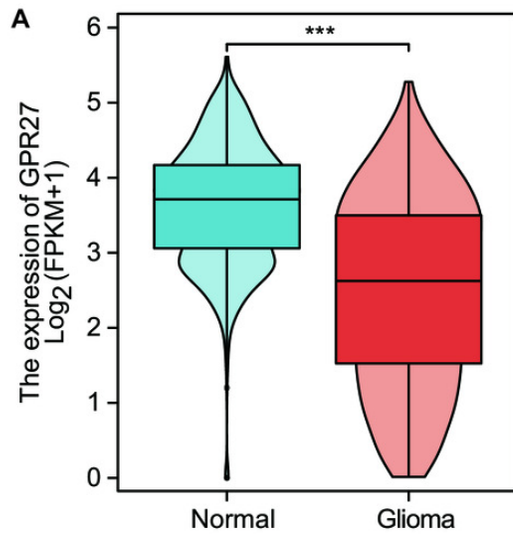
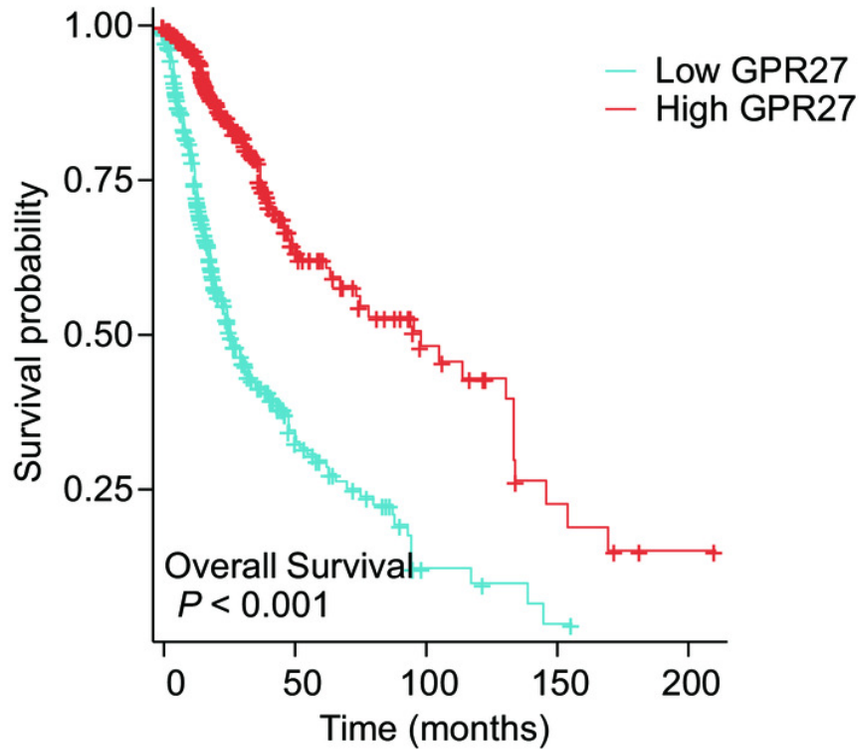


Figure 2

Figure 2. Prognostic significance of GPR27 on glioma patients.

A) Kaplan-Meier survival analysis of glioma patients based on GPR27 expression level. Patients with lower GPR27 level had significantly worse prognosis compared to those with higher GPR27 level ($P < 0.001$). B) A nomogram established based on the multivariate survival analysis. The variables in the nomogram included patients' age, gender, WHO grade, IDH status, 1p/19q co-deletion, and GPR27 expression level.

A



B

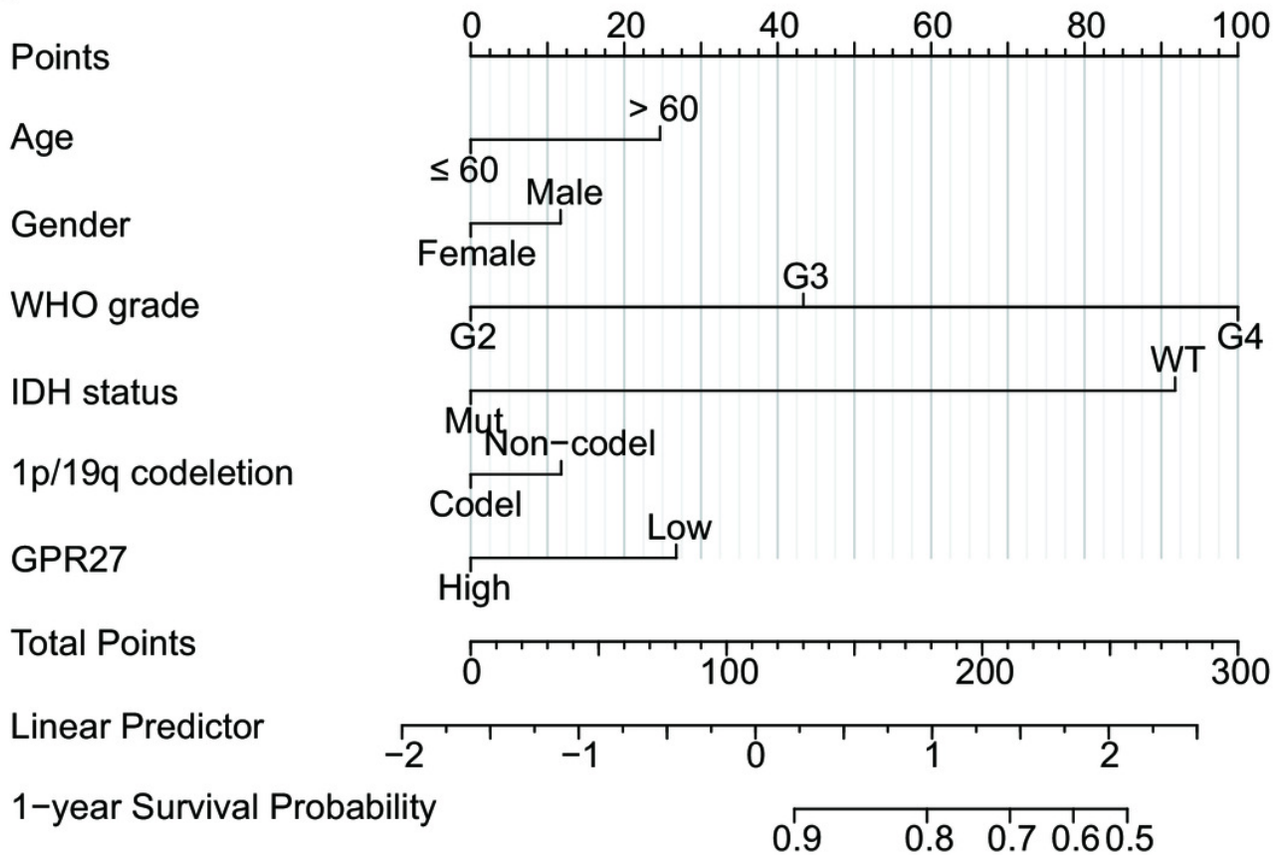


Figure 3

Figure 3. GPR27-knockdown inhibit glioma progression.

A) Western blot analysis confirmed the successful knockdown of GPR27 expression in U87 and U251 cell lines. B) CCK-8 assay results indicated that silencing GPR27 significantly inhibited the proliferation of U87 cells compared to the control group. C) CCK-8 assay results indicated that silencing GPR27 significantly inhibited the proliferation of U251 cells compared to the control group.

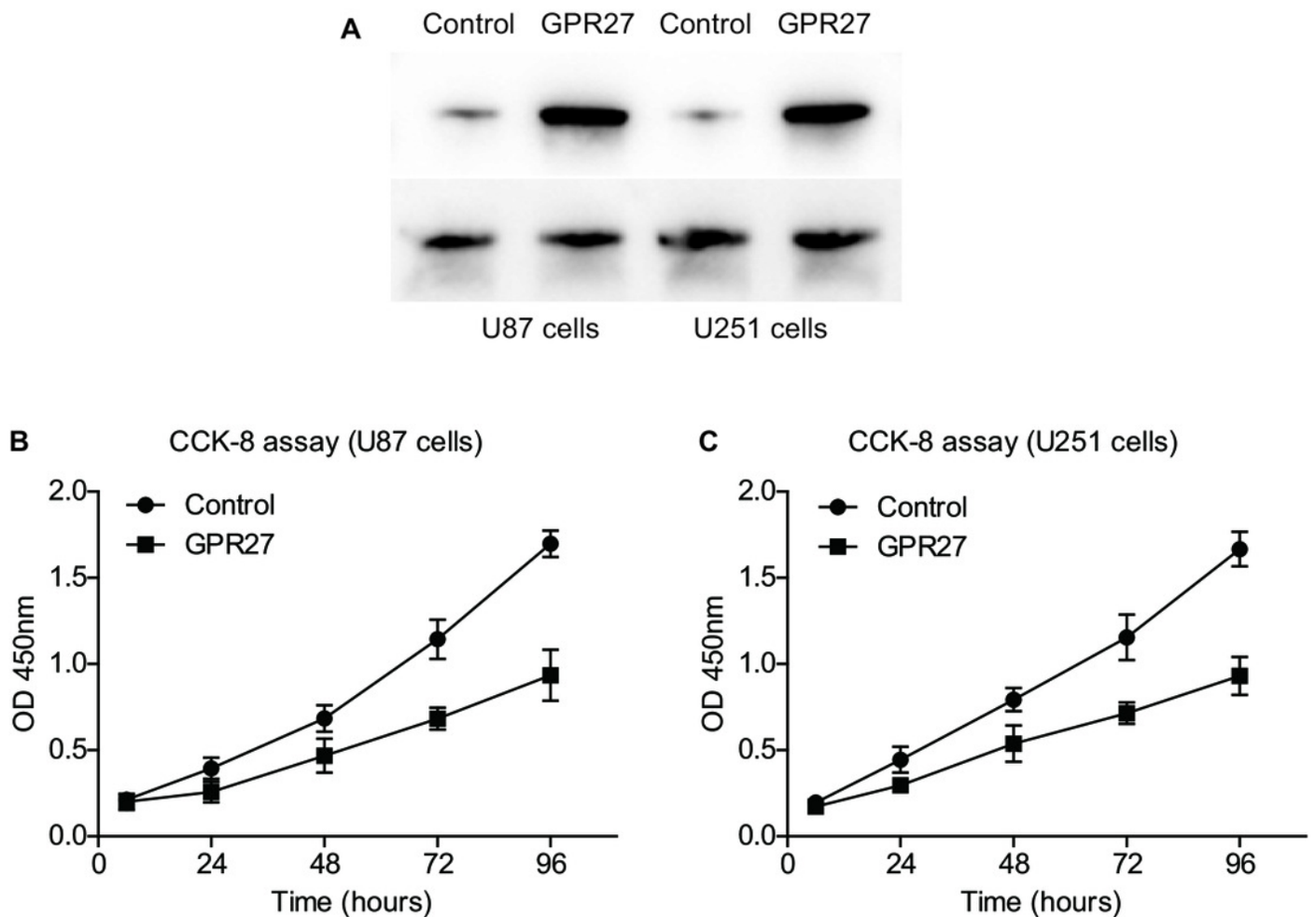


Figure 4

Figure 4. The correlation between immune cell infiltration and GPR27 expression in glioma.

A) Spearman analysis result showed the correlation between the infiltration of 24 types of immune cells and GPR27 expression in glioma tissues. B) and C) showed representative infiltration level of pDC and macrophages in cells with different GPR27 level. DC, dendritic cells; pDC, plasmacytoid DC; aDC, activated DC; iDCs, immature DCs; NK, natural killer cells; Tgd, T gamma delta; TReg, regulatory T cells; Tem, T effector memory; Tcm, T central memory; Th1 cells, type 1 Th cells; Th2 cells, type 2 Th cells; Th17 cells, type 17 Th cells; TFH, T follicular helper.

