XuefuZhuyu Tang exerts antitumor effects by inhibiting glioma cell metastasis and invasion via regulating tumor microenvironment

Background: XuefuZhuyu Tang (XZT) is a traditional Chinese herb used for destagnation and is currently being used for oncotherapy. This study was intended to assess the effects of XZT on glioma along with its anticancer mechanism.

Materials and methods: U251 cells were divided into five groups: CNC (cells were cultured with normal saline), TSC (cells were treated with TaohongSiwu Tang [TST]), XSC (cells were treated with XZT), THC (cells were treated with homogenate of TST), and XHC (cells were treated with homogenate of XZT). The mRNA and protein expression of VEGF/VEGFR, CXCR4/CXCL12, and TIMP1/MMP9/MMP2 were measured by reverse transcription-polymerase chain reaction (RT-PCR) and Western blotting, respectively. Moreover, MTT assay, transwell assay, wound-healing assay, and flow cytometry were conducted to assess the cell viability, cell migration and invasion, cell motility, and cell apoptosis of U251 cells, respectively. In vivo, three mice models (group CNM, gavaging saline; group TSM, gavaging TST; group XZM, gavaging XZT) were constructed after establishing xenograft mice models. Then, models were examined using hematoxylin and eosin staining, RT-PCR, and Western blotting.

Results: In vitro, XZT significantly upregulated TIMP1 expression and downregulated the expression of VEGF, VEGFR, CXCR4, CXCL12, MMP9, and MMP2 in U251 cells (all \(P<0.05\)). In addition, XZT inhibited cell proliferation, invasion, and migration and induced cell apoptosis. In vivo, the average expression level of VEGF, CXCL12, MMP9, and MMP2 was downregulated in the XZM group compared with the control and TSM groups (all \(P<0.05\)). Tumor volumes in the XZM group were significantly lower than those in the CNM and TSM groups (all \(P<0.05\)).

Conclusion: XZT may suppress glioma growth and decrease expression levels of VEGF, CXCL12, MMP9, and MMP2. We speculate that XZT may be a potential therapeutic herb for curing glioma.

Keywords: XuefuZhuyu Tang, glioma, tumor microenvironment, VEGF, CXCL12, TIMP1, MMP9, MMP2

Introduction

Glioma is commonly discovered in the central nervous system and primarily occurs in the brain and glial tissues. Glioma accounts for ~81% of malignant brain tumors with unexpectedly high mortality, and its incidence, though very rare, ranges from 4.67 to 5.73 per 100,000 persons. It is reported that certain environmental carcinogens, occupations, and diet (N-nitroso compounds) may increase the risk of glioma. Current treatments for glioma include surgery and concomitant temozolomide chemotheraphy. The presence of blood–brain barrier, which prevents macromolecules from entering...
into the brain, limits the efficacy of immunotherapeutic approaches and gene therapy.\textsuperscript{3} Although it has been suspected that tumor growth and progression is driven by the tumor itself, recent studies have suggested that tumor microenvironment (TME) is also associated with tumor progression and metastasis.\textsuperscript{3} TME is a rich and robust stromal cell matrix, which is composed of fibroblasts, endothelial cells, pericytes, leukocytes, and extracellular matrix. Moreover, TEM plays a decisive role in tumor growth and may lead us to develop groundbreaking therapeutic targets for tumors.\textsuperscript{4}

Growing studies have attempted to unfold the underlying mechanism between TME and tumor since TME has been acknowledged as a causal factor for the development of tumors. As receptors of CXCL12, both CXCR4 and CXCR7 have been discovered to be highly expressed in tumor cells, which indirectly proved the interplay among stem-like cancer cells, differentiated glioma cells, and TME. Particularly, the circulation is completed via secreted chemokines (eg, CXCL12), which recruit various kinds of molecules, including fibroblasts, endothelial, mesenchymal, and inflammatory cells through CXCR4 to tumor tissues.\textsuperscript{5} Apart from CXCL12, VEGF is believed to be a key growth factor that is responsible for neovasculature; therefore, aberrantly elevated VEGF along with florid vascularization has been considered as a common characteristic of glioma. Meanwhile, it has been suggested that VEGF could regulate tumor invasion by recruiting the protein to a heterocomplex, including VEGFR2, which further suppresses tumor migration.\textsuperscript{6}

Recently, antiangiogenic therapies targeting VEGF and its receptors have been developed to inhibit glioma growth. However, the clinical outcomes are transient, and recurrence is usually observed.\textsuperscript{7} MMPs are zinc-dependent endoproteases that are able to degrade the extracellular matrix and mediate tumor growth and metastasis. The tissue inhibitors of metalloproteinases (TIMPs) primarily prevent the matrix from degradation through the inhibition of MMPs.\textsuperscript{8}

Traditional Chinese medicine (TCM) has been widely used and practiced for thousands of years. \textit{TaohongSiwu Tang (TST)} and \textit{XiefuZhuyu Tang (XZT)} are famous traditional Chinese herbal medicines. XZT includes the following eleven kinds of herbs: Sichuan lovage root (chuanxiong), red peony root (chishao), rehmannia root (shengdi), platycodon root (jiegeng), peach seed (taoren), safflower (honghua), hare’s ear root (chailiu), Chinese angelica (danggui), two-toothed achyranthes root (niuxi), orange fruit (zhiquiao), and liquorice. The formulation of XZT is well known for Qi circulation promotion and blood stasis removal.\textsuperscript{9} Many herbs in the medicine belong to plant functional foods and play protective roles in cancer prevention/treatment.\textsuperscript{10,11} Since tetramethylpyrazine (TMP) extracted from the Chinese herb chuanxiong has positive effects on tumor care, XZT is considered as a treatment for cardiovascular and chronic liver diseases.\textsuperscript{9,12} The effects of XZT on fibrotic liver include not only the inhibition of collagen deposition but also the antiangiogenesis.\textsuperscript{12}

Though a large number of studies have revealed the antiangiogenic effects of XZT, the molecular mechanisms of XZT that are related to glioma are still ambiguous. As a result of this, we used both in vitro and in vivo models to study the effect of XZT on tumor cells and expressions of VEGF, CXCL12, MMP9, and MMP2. We suspected that XZT inhibits glioma development and metastasis by regulating the extracellular microenvironment of glioma cells, which may further provide new insights for glioma therapies.

**Materials and methods**

**Animals, cells, and groups**

Herbs of TST and XZT used in this study were purchased from The First Affiliated Hospital of Guangzhou University (Guangzhou, People’s Republic of China). Then dry herbs were dissolved in sterile 0.9% NaCl to form appropriate concentrations for usage. Normal saline was applied as the control in all experiments.

Sixteen male SD rats with an average weight of ~120 g and 24 male BALB/c nude mice with an average weight of ~10 g were purchased from the Laboratory Animal Center of Southern Medical University (Guangzhou, People’s Republic of China). Animals were housed in a specific pathogen-free environment. The relative humidity and temperature were set at 50%±10% and 25°C±1°C, respectively. Animals were subjected to 10 hour light and 14 hour dark cycles per 24 hours.

U251 glioma cell lines were offered by the Institute of Biochemistry and Cell Biology (Shanghai, People’s Republic of China). Cells were grown at 37°C in Roswell Park Memorial Institute 1640 medium with 10% fetal bovine serum, 100 U/mL penicillin, and 100 mg/L streptomycin in a humidified atmosphere containing 5% CO\textsubscript{2}.

Cultured cells were divided into the following five groups: group CNC (U251 cells cultured with normal saline, control group), group TSC (U251 cells cultured with serum of TST-treated SD rats, medicine control group), group XSC (U251 cells cultured with serum of XZT-treated SD rats), group THC (U251 cells cultured with homogenate of TST-treated SD rats), and group XHC (U251 cells cultured with homogenate of XZT-treated SD rats). The above experiments
were carried out for 1 week. This study and all study protocols were approved by the ethics committee of The First Affiliated Hospital of Guangzhou University of Traditional Chinese Medicine, and this study complied with the Guidelines for Animal Experimentation Committee at Guangzhou University of Traditional Chinese Medicine.

**RNA extract and real-time reverse transcription-polymerase chain reaction**

The mRNAs of CXCR4, CXCL12, VEGF, VEGFR, TIMP1, MMP9, and MMP2 were taken from cells and tissues using the miRNeasy Mini Kit (Qiagen NV, Venlo, the Netherlands) under the manufacturer’s instructions. Reverse transcription-polymerase chain reaction (RT-PCR) was performed to obtain cDNA template using the ReverTra Ace qPCR RT Kit (Toyobo, Osaka, Japan) along with the CFX96 Touch RT-PCR Detection System (Bio-Rad Laboratories Inc., Hercules, CA, USA). The primer sequences are shown in Table 1. Relative expression quantity of target genes was normalized to that of the internal control group (β-actin) and was calculated using the $2^{-\Delta\Delta CT}$ method. The experimental procedures were repeated for at least three times.

**Western blotting**

Western blotting was used to detect the expression level of targeted proteins. Cellular extracts were lysed by the lysis buffer used in radioimmunoprecipitation assay. Supernatant was collected after centrifuging and lysing the cellular extracts at 12,000 rpm for 15 minutes at 4°C. Cell proteins were isolated using sodium dodecyl sulfate–polyacrylamide gels and were transferred to polyvinylidenedifluoride membranes (Bio-Rad Laboratories Inc.). Then membranes with isolated cell proteins were blocked for 1 hour and detected using primary antibodies, including anti-CXCR4 (1:100), anti-CXCL12 (1:100), anti-VEGF (1:500), anti-VEGFR (1:500), anti-TIMP1 (1:300), anti-MMP9 (1:300), anti-MMP2 (1:300), and anti-β-actin (1:100) (Santa Cruz Biotechnology Inc., Dallas, TX, USA) at 4°C overnight. Membranes were further washed thrice by $\times 1$ phosphate-buffered saline (PBS) Tween-20 (10 min/time). After that, horseradish peroxidase-conjugated secondary antibodies were added to membranes, which were incubated for another 1 hour. Finally, membranes were washed thrice again using Tris buffered saline Tween (10 min/time). The immunobinding signals were detected by the chemiluminescence method (KeyGen Biotech Co. Ltd, Nanjing, People’s Republic of China). Relative protein expression was subjected to densitometry analysis using the Image-Pro Plus Version 6.0 software (Media Cybernetics., Inc, Washington, IL, USA) and calculated according to the β-actin reference bands.

**MTT assay**

MTT (3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2-H-tetrazolium bromide) assay was carried out to assess the viability of U251 cells. U251 glioma cells were incubated for 4 hours at 37°C in a 96-well plate, and then 0.5% DMSO was added to each well in the plate. Absorbance was detected by a fluorescence plate reader at 560 nm (Bio-Rad Laboratories Inc.). Cell viability was determined by the optical density ratio of a treated group over an untreated control group. The experiments were repeated for three times.

### Table 1 Primer sequences of targeted genes

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer</th>
<th>Size (bp)</th>
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| CXCR4 | For 5'-ACTCGGTGCTCGGAGATGATG-3'  
          Rev 5'-GTGGTACGCTTTTCTGTGATG-3' | 558       |
| CXCL12| For 5'-GTCGCGCAATGGACCTC-3'  
          Rev 5'-CTTGGGAAAGCCTCCTTTA-3' | 1,180     |
| VEGF  | For 5'-TGTGCTCTCCCTTCCACCAT-3' 
          Rev 5'-TGTGCTCTCCCTTCCCATAG-3' | 64        |
| VEGFR | For 5'-TGAAAGGAAAGCCAGATGCTCAT-3' 
          Rev 5'-CAGGTATATTGGCTTCCTCAC-3' | 111       |
| TIMP1 | For 5'-TGACGATTGGCAGCTC-3'  
          Rev 5'-GGTACCGTGGCAGGAGG-3' | 551       |
| MMP9  | For 5'-GTTGCCGAGTGAATTGA-3'  
          Rev 5'-TTTACATGGCAGCAAGC-3' | 109       |
| MMP2  | For 5'-GCCTCCGCGGTAG-3'  
          Rev 5'-GAGGCACTTGCCAAAGG-3' | 133       |
| β-Actin| For 5'-TCGACACAGGGCTTTCCCATCAT-3' 
          Rev 5'-AAGGTGTTGGGTCCAGATTTC-3' | 241       |

**Abbreviations:** For, forward; Rev, reverse.
Transwell assay
A transwell chamber (8 mm pore size; Costar, High Wycombe, UK) was placed into the wells of 12-well culture plates in order to measure cell migration and invasion. A total of 5×10⁴ cells were added to the upper chamber and cultured with serum-free medium, while the lower chamber was filled with the complete medium. Cells in the upper chamber were carefully removed by a cotton swab after 24-hour incubation, while cells migrated or invaded into the lower surface were fixed and stained with eosin. Matrigel was specifically added to the medium for cell invasion assay. The number of migrated or invaded cells was counted for five random fields under the microscope (Olympus Corporation, Tokyo, Japan), and three independent assays were conducted.

Wound-healing assay
The wound-healing assay was conducted to assess cell motility. U251 cells in each group were seeded into 12-well plates 1 day prior to scratching. A scratching wound was performed by scraping the middle of the cell monolayer using a sterile micropipette tip. After removing the detached cells by PBS, cells were cultured with complete medium. Image of cells migrating into the wound was captured at 0 hour and 24 hours using a microscope recording the distance of U251 cell migration.

Cell apoptosis analysis
Apoptosis was evaluated using the Annexin V-FITC Apoptosis Detection Kit. U251 cells were seeded in six-well plates, incubated overnight, and then treated with serum (TST or XZT) and homogenate (within TST or XZT) for 3 days. Cells were harvested by trypsinization, washed with PBS, and finally centrifuged. Cell pellet was resuspended in the binding buffer and incubated with Annexin V-fluorescein isothiocyanate (FITC). After that, cells were again centrifuged, washed with PBS, and resuspended in the binding buffer containing PI solution in the dark. Finally, cells were analyzed by the approach of flow cytometry (Beckman FC400 MPL; Beckman Coulter Inc., California, IL, USA). Each experiment was independently carried out in triplicate.

Xenograft mouse model
U251 cells were harvested, washed with PBS, and resuspended in a mixture of culture medium and Matrigel (BD Biosciences, San Jose, CA, USA). Then, U251 cells (1×10⁶ per site) were subcutaneously implanted into the two flanks of all nude mice.

A total of 24 mice were randomly allocated into the following three groups with equal size when tumors grew to the level of ~200 mg: group CNM (gavaging saline), group TSM (gavaging TST), and group XZM (gavaging XZT). Treatments were administered once a day for the period of 28 days, and the concentrations of TST and XZT were both set at 100 µM.

Tumor size in the two flanks was measured twice per week using calipers (tumor volume = [length × width³] × 0.5). Tumor growth was monitored twice per week for the period of 30 days. Mice were anesthetized and sacrificed at the end of the experiment, and tumors were extracted from mice for imaging and weighing. Tumors were fixed with 4% paraformaldehyde, dehydrated, and embedded with paraffin. Then, 5 mm sections were cut from tumor tissues and stained with hematoxylin and eosin and observed using a microscope. The experimental procedures were complied with the Animal Management Rule of the Chinese Ministry of Health (Documentation 55, 2001), and the experimental protocol was approved by the ethics committee of The First Affiliated Hospital of Guangzhou University of Traditional Chinese Medicine.

Statistical analysis
All statistical analyses were performed by SPSS 18.0 software (SPSS Inc., Chicago, IL, USA). Continuous data were expressed in the form of mean ± SD. The two-tailed Student’s t-test or one-way analysis of variance was used to assess between-group comparisons, and P<0.05 provided evidence for statistical significance.

Results
XZT affects expressions of VEGF/VEGFR, CXCR4/CXCL12, TIMP1/MMP9/MMP2 in U251 glioma cells in vitro
Both RT-PCR and Western blotting were conducted to assess whether XZT is able to affect the expression of VEGF/VEGFR, CXCR4/CXCL12, and TIMP1/MMP9/MMP2 in glioma U251 cells. As shown in Figure 1, the TIMP1 mRNA expression level in U251 cells treated with XZT was significantly higher than that in U251 cells treated with saline or TST, while the mRNA expression levels of VEGF, VEGFR, CXCR4, CXCL12, MMP9, and MMP2 exhibited the opposite trend (P<0.05). Similarly, Western blotting results indicated that TIMP1 in U251 cells treated with XZT expressed significantly higher than that in U251 cells treated with saline or TST, while the expressions of VEGF, VEGFR, CXCR4, CXCL12, MMP9, and MMP2 in U251 cells treated with XZT were significantly lower than those in U251 cells treated with saline or TST (P<0.05) (Figure 2). Our data revealed that XZT significantly upregulated the expression of TIMP1 and...
downregulated the expression of VEGF, VEGFR, CXCR4, CXCL12, MMP9, and MMP2 in U251 glioma cells in vitro.

**XZT inhibits tumor cell proliferation of U251 glioma cell lines in vitro**

Furthermore, the MTT assay was performed to evaluate the viability of U251 glioma cells treated with XZT, TST, and saline. Briefly, cells were treated with five different treatment groups for a period of 24 hours, 48 hours, and 72 hours. As suggested by Figure 3, XZT inhibited U251 glioma cell proliferation in a time-dependent manner. Moreover, the inhibition rate in the XSC group was significantly higher than that in the CNC and TSC groups when cells were treated for a period of 48 hours or 72 hours (all \( P<0.05 \)).

**XZT inhibits glioma cell invasion and migration**

As suggested by Figure 4, the number of migrated cells in group XSC (80.12±4.13) was significantly less than that in group CNC (189.58±10.05) or group TSC (149.68±5.16) (all \( P<0.05 \)) after treatments have been applied for a period of 7 days, suggesting that XZT serum can effectively inhibit U251 glioma cell migration. Besides that, XZT homogenate served as a glioma cell migration inhibitor. The number of migrated cells in group XHC (79.78±6.20) was significantly less than those in group CNC (189.58±10.05) and group THC (166.44±15.02) (all \( P<0.05 \), Figure 4). As shown in Figure 5, results from wound-healing assays illustrated that group XSC suppressed cell migration after treatments have been applied for 12 hours, and the same trend was observed after treatments have been applied for 24 hours.

**XZT induces apoptosis of U251 glioma cells in vitro**

Flow cytometry with Annexin V-FITC/PI staining was used to analyze whether XZT was able to induce U251 glioma cell apoptosis. As suggested by Figure 6A–C, the apoptosis rate in group XSC was significantly higher compared to that in group CNC or TSC (all \( P<0.05 \)), which indicated that XZT serum can greatly promote U251 glioma cell apoptosis. XZT
**Abbreviations:** sD, sprague Dawley; CNC, control group; TSC, XSC, XHC, medicine control group. group Xsc: U251 cells were cultured with serum of sD rats treated with homogenate of sD rats treated with homogenate of sD rats treated with homogenate of sD rats treated with normal saline, control group. group Tsc: U251 cells were cultured with serum of sD rats treated with homogenate of sD rats treated with normal saline, control group. group Xhc: U251 cells were cultured with serum of sD rats treated with homogenate of TSC. group Thc: U251 cells were cultured with serum of SD rats treated with homogenate of TST. Group XHC: U251 cells were cultured with serum of SD rats treated with homogenate of TST.

**Notes:** *Versus group CNC, P<0.05; *Group XSC versus group TSC and group XHC versus group THC, P<0.05. Group CNC: U251 cells were cultured with normal saline, control group. Group TSC: U251 cells were cultured with serum of SD rats treated with TST, medicine control group. Group XSC: U251 cells were cultured with serum of SD rats treated with XZT. Group THC: U251 cells were cultured with serum of SD rats treated with homogenate of TST. Group XHC: U251 cells were cultured with serum of SD rats treated with homogenate of XZT.

**Figure 2** The expressions of targeted proteins in U251 cells with different treatments by Western blotting.

**Notes:** β-Actin was used as control. Group CNC: U251 cells were cultured with normal saline, control group. Group TSC: U251 cells were cultured with serum of SD rats treated with TST, medicine control group. Group XSC: U251 cells were cultured with serum of SD rats treated with XZT. Group THC: U251 cells were cultured with serum of SD rats treated with homogenate of TST. Group XHC: U251 cells were cultured with serum of SD rats treated with homogenate of XZT.

**Abbreviations:** SD, Sprague Dawley; TST, TaohongSiwu Tang; XZT, XuefuZhuyu Tang.

Figure 3 XZT inhibits U251 glioma cell proliferation detected by the MTT assay.

**Notes:** The five groups of cells were cultured for 24 hours, 48 hours, and 72 hours. Results are expressed relative to the cell density of group CNC. *Versus group CNC, P<0.05; *Group XSC versus group TSC and group XHC versus group THC, P<0.05. Group CNC: U251 cells were cultured with normal saline, control group. Group TSC: U251 cells were cultured with serum of SD rats treated with TST, medicine control group. Group XSC: U251 cells were cultured with serum of SD rats treated with XZT. Group THC: U251 cells were cultured with serum of SD rats treated with homogenate of TST. Group XHC: U251 cells were cultured with serum of SD rats treated with homogenate of XZT.

**Abbreviations:** MTT, 3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide; SD, Sprague Dawley; TST, TaohongSiwu Tang; XZT, XuefuZhuyu Tang.

Figure 4 The invasion and migration ability of U251 glioma cells at 7 days after treatment was detected by transwell assay.

**Notes:** *Versus group CNC, P<0.05; *Group XSC versus group TSC and group XHC versus group THC, P<0.05. Group CNC: U251 cells were cultured with normal saline, control group. Group TSC: U251 cells were cultured with serum of SD rats treated with TST, medicine control group. Group XSC: U251 cells were cultured with serum of SD rats treated with XZT. Group THC: U251 cells were cultured with serum of SD rats treated with homogenate of TST. Group XHC: U251 cells were cultured with serum of SD rats treated with homogenate of XZT.

**Abbreviations:** SD, Sprague Dawley; TST, TaohongSiwu Tang; XZT, XuefuZhuyu Tang.

Figure 5 The scratch repaired ability of U251 glioma cells at 12 hours and 24 hours after treatment was detected by wound-healing assay.

**Notes:** Group XSC was compared with group CNC and group TSC, and group XHC was compared with group CNC and group THC. Group CNC: U251 cells were cultured with normal saline, control group. Group TSC: U251 cells were cultured with serum of SD rats treated with TST, medicine control group. Group XSC: U251 cells were cultured with serum of SD rats treated with XZT. Group THC: U251 cells were cultured with serum of SD rats treated with homogenate of TST. Group XHC: U251 cells were cultured with serum of SD rats treated with homogenate of XZT.

**Abbreviations:** SD, Sprague Dawley; TST, TaohongSiwu Tang; XZT, XuefuZhuyu Tang.

**XZT suppresses glioma cell proliferation in vivo**

Volumes of each xenograft tumor were measured after they were extracted from the xenograft mice. As displayed...
in Figure 7A and B, tumor volumes in group XZM (542.25±84.51 mm³) were significantly lower than those in group CNM (1,230.33±164 mm³) and group TSM (1,052.63±156.89 mm³) (all P<0.05) after treatments had been applied for a period of 30 days. Moreover, we conducted haematoxylin and eosin stain to observe the morphology of tumors (Figure 7C). There was no significant abnormality observed in tumor cells of group CNM. However, tumor cells in group TSM and group XZM migrated into the brain to some extent, particularly glioma cells in group XZM showed a higher migration pattern (Figure 7C), indicating that glioma cells treated with XZT for a period of 30 days may be associated with a stronger suppression of cell proliferation compared to that in group CNM or group TSM. As suggested by Figure 7D, RT-PCR and Western blotting in vitro revealed that expressions of VEGF, CXCL12, MMP9, and MMP2 were downregulated in XZT group in comparison to the control and TST groups.

Discussion
As a common brain neoplasm, glioma has poor prognosis due to the lack of effective treatments.\(^\text{13}\) So far, the median survival time of patients with malignant glioma is ~1 year.\(^\text{15}\) A variety of studies both in vivo and in vitro have demonstrated that TCM, such as Chinese herbs, may be considered as alternative treatments for cancer.\(^\text{14,15}\) For instance, TMP extracted from chuanxiong, which is a natural Chinese herb, has been suggested as a potential therapy for glioma due to its neural protection actions.\(^\text{16–18}\)

XZT is a widely known TCM formulation consisting of eleven herbal substances, including chuanxiong. Our findings indicated that XZT exerted its anticancer function by inhibiting the proliferation, invasion, and migration of glioma cells and inducing U251 glioma cell apoptosis in vitro. Similar results reported that the mean survival time and life span of mice suffering from experimental liver tumors were notably improved by XZT treatment compared to the control group.\(^\text{19}\) Therefore, XZT may have the potential to be a new therapeutic agent for patients with glioma.

XZT is a typical formulation for invigorating blood circulation and dissipating blood stasis and was first described by Yi Lin Gai Cuo, which was a monograph on TCM cases in Qing Dynasty in AD 1830.\(^\text{20}\) Previous studies have provided evidence that XZT is able to treat cardiovascular and liver diseases. For example, a study conducted by Zhou et al\(^\text{12}\) suggested that XZT could not only restrain collagen deposition but also exert its antiangiogenic effect on fibrotic liver through relieving hypoxia and preserving liver sinusoidal endothelial cell functions, thus restraining...
Figure 7 XZT inhibits the glioma tumor growth at 30 days after gavage in vivo.

Notes: (A) The representative photo of glioma tumors in each xenograft model group. (B) The tumor volumes decreased significantly with the group XZM compared to the group CNM or group TSM. *Versus group CNM, P < 0.05; #Group XZM versus group TSM, P < 0.05. (C) The number was decreased significantly in the group XZM compared to the group CNM or group TSM analyzed by H&E stain. Magnification ×200. (D) Relative expressions of MMP2, MMP9, VEGF, and CXCL12 were downregulated in xenograft mice of the group XZM compared to those of the group CNM or group TSM. *Versus group CNM, P < 0.05; #Group XZM versus group TSM, P < 0.05. Group CNM, gavaging saline; group TSM, gavaging TST; group XZM, gavaging XZT.

Abbreviations: H&E, hematoxylin and eosin; TST, TaohongSiwu Tang; XZT, XuefuZhuyu Tang.

Liver fibrosis. Moreover, Hwang et al21 showed that XZT could significantly affect the progression of syndromes and inflammatory markers observed in patients with unstable angina pectoris after treating with percutaneous coronary intervention.

This study enabled us to discover that XZT remarkably upregulated the expression of TIMP1 and significantly downregulated expression of VEGF, VEGFR, CXCR4, CXCL12, MMP9, and MMP2 in U251 glioma cells in vitro, implying that XZT may affect glioma cells via regulating TME.
Similarly, our investigations in vivo revealed that tumor volumes in group XZM were significantly lower than those in group CNM and group TSM. Moreover, the expressions of VEGF, CXCL12, MMP9, and MMP2 were downregulated in group XZM compared with group CNM and group TSM, which was consistent with the results obtained from RT-PCR and Western blotting in vitro. Our findings of CXCR4/CXCL12 pathway were agreed with those by Chen et al, which suggested that downregulated expression of chemokine receptor CXCR4 by TMP suppressed the activities of glioma cells, including proliferation, migration, and invasion, and reduced the release of glutamate from cerebral neurocytes. These results indicated that the CXCR4/CXCL12 pathway may be considered as a new and potential mechanism in which TMP-mediated tumor is depressed and neural functions are preserved.16

To date, growing studies have supported that TME is an emerging therapeutic target for glioma. For instance, Yang et al22 reported that the activation of CXCR4 could induce glioma growth in which c-AMP is accumulated in glioma cells, whereas the specific CXCR4 antagonist AMD 3465 is able to inhibit tumor growth both in vivo and in vitro, indicating that the CXCR4/CXCL12 axis may possess critical roles in tumor progression. Yang et al also demonstrated that high-LET carbon ion radiation was preferred to low-LET X-rays with respect to suppressing the migration of glioma cells. Furthermore, their study showed that X-ray radiation enhanced the secretion of VEGF, MMP2, and MMP9, which can further degrade basement membrane and extracellular matrixes when tumor progression occurs.23–26 Qazi et al27 also demonstrated that fluid shear stress may decrease the motility of glioma cells through modulating MMPs’ activation and expression. Furthermore, Najbauer et al noted that VEGF could contribute to the generation of distended and glomerulus-like vessels in glioma cells after treating with MMPs.28 However, it is still difficult to explain whether inhibitory effects of XZT on VEGF/VEGFR, CXCR4/CXCL12, and TIMP1/MMP9/MMP2 pathway are independent or coadjusted. Therefore, further research is required to clarify the relationship between VEGF/VEGFR, CXCR4/CXCL12, and TIMP1/MMP9/MMP2 in the inhibitory mechanism of XZT.

Conclusion

Our study suggested that XZT could possess many properties, including the inhibition of glioma cell proliferation, protection of neurons from excitotoxicity induced by glioma cells, and improving blood–brain barrier penetration. Apart from that, XZT may suppress the proliferation, invasion, migration, and growth of glioma cells through regulating the expression levels of microenvironment relative factors. Importantly, multilevels, multtargets, and multiple pathways were observed in the anticancer mechanism of XZT. These significant findings may provide a profound explanation for the molecular basis of XZT in treating glioma and may direct future research, which will be able to clarify the clinical benefits of XZT.

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Disclosure

The authors report no conflicts of interest in this work.

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