siRNA targeting stathmin inhibits invasion and enhances chemotherapy sensitivity of stem cells derived from glioma cell lines

Yuwen Song¹, Luyan Mu¹, Xuezhe Han², Xiaoqian Liu¹*, and Songbin Fu³

¹Department of Neurosurgery, the Fourth Affiliated Hospital of Harbin Medical University, Harbin 150001, China
²Neurosurgery and Vascular Biology Program, Children’s Hospital Boston/Harvard Medical School, Boston, MA 02115, USA
³Department of Genetics, Harbin Medical University, Harbin 150081, China

*Correspondence address: Tel/Fax: + 86-451-82576656; E-mail: lxq1960@hotmail.com

Glioma is one of the most highly angiogenic tumors, and glioma stem cells (GSCs) are responsible for resistance to chemotherapy and radiotherapy, as well as recurrence after operation. Stathmin is substantial for mitosis and plays an important role in proliferation and migration of glioma-derived endothelial cells. However, the relationship between stathmin and GSCs is incompletely understood. Here we isolated GSCs from glioma cell lines U87MG and U251, and then used siRNA targeting stathmin for silencing. We showed that silencing of stathmin suppressed the proliferation, increased the apoptosis rate, and arrested the cell cycle at G2/M phase in GSCs. Silencing of stathmin in GSCs also resulted in inhibited the migration/invasion as well as the capability of vasculogenic mimicry. The susceptibility of GSCs to temozolomide was also enhanced by stathmin silencing. Our findings suggest stathmin as a potential target in GSCs for glioma treatment.

Keywords stathmin; invasion; temozolomide; glioma stem cells

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Introduction

Glioma is the most common and malignant primary brain tumors in adults, which has a poor prognosis and displays unique biological features especially in the network of neoplastic blood vessels, invasion, and metastasis. The median survival time of patients with glioblastoma is still only 1 year despite positive surgery and therapy [1,2]. Tumor microvessel endothelial cells have been shown to be morphologically different from normal endothelial cells, with elevated migration and resistance to necrosis [3]. Thus, anti-angiogenesis therapy is important for glioblastoma multiforme (GBM) treatment. Previous studies have identified a small population of tumor cells called glioma stem cells (GSCs). GSCs were first isolated in 2003 [4] and were responsible for GBM initiation, propagation, chemical therapy resistance, and glioma recurrence [5,6]. How to inhibit GSCs becomes the hot topic of glioma research.

Recently, a number of researches have demonstrated that vasculogenic mimicry (VM) plays a central role in the vascularization of GBM [7–9]. VM was an alternative vascular mechanism, which was first described and named by Maniotis et al. [10] in 1999. VM describes the ability of aggressive tumor cells to form vasculogenic-like networks which were associated with their high plasticity. VM is also involved in more aggressive tumor biology and can increase tumor-related mortality [11,12]. GSCs, which are the response for malignance of GBM and capable of trans-differentiation into vascular nonendothelial cells, have a strong ability of VM [13,14].

Stathmin, also known as oncoprotein 18 (OP18), is a member of the microtubule destabilizing protein family. It regulates microtubule dynamics during cell-cycle progression [15]. Most studies have demonstrated that the expression of stathmin is associated with tumor progression and unfavorable long-term prognosis. We previously reported that stathmin 1 is over-expressed in human glioma, and the inhibition of stathmin expression in high-grade glioma-derived endothelial cells significantly inhibits cell proliferation, migration, and invasion [16].

In this study, we isolated GSCs from glioma cell lines and investigated whether stathmin 1, which is extensively researched in the stathmin family, could also affect the proliferation and invasion ability of GSCs, as well as chemotherapy sensitivity to temozolomide (TMZ) to provide a potential target for anti-angiogenic treatment of glioma.

Materials and Methods

Cell culture and isolation of GSCs

Malignant glioma cell lines U87MG and U251 were obtained from American Type Culture Collection (Rockville, USA)
and cultured in complete Dulbecco’s modified Eagle’s medium (DMEM; Invitrogen Tech, Shanghai, China). Then glioma stem-like cells were successfully obtained from U87MG and U251 cell lines following the procedure described previously [4]. Identification of GSCs was carried out by western blot analysis using antibodies against stem cell markers (Oct3/4, MDR1, Sox2 and Nestin; Abcam, Cambridge, UK).

RNA interference
The Stealth siRNA against stathmin was designed and synthesized by Invitrogen Tech. The sequences were 5’-GCUUC UUCUGAUACCAGGUGAAAG-3’, 5’-CUUUCACCU GGAAUACAGAAGAAGC-3’, and 5’-GAGCUGAUUCU CAGCCUCGGUCAA-3’. GSCs grown to a density of 1 × 10^5 cells/ml were transfected in duplicate with 80 nM of a siRNA pool (three siRNA duplexes) targeting stathmin. The mixture of 4 μl siRNAs and 6 μl lipofectamine 2000 (Invitrogen Tech) was added onto the cells after 15-min incubation at room temperature. After overnight incubation, the cells were switched to complete DMEM for 2 days. In this experiment, scramble sequence was applied as the control group.

GSC treated with TMZ
TMZ obtained from Sigma-Aldrich Co. LLC (St Louis, USA) was dissolved in DMSO to make the stock solution at 10 mM which was then diluted into gradient concentrations. Then GSCs were cultured in 96-well plates and treated with gradient TMZ. Then LD50 of TMZ on GSCs was calculated, and cell apoptosis analysis and cell cycle assay were also performed.

Cell proliferation assay
GSCs were cultured in 96-well plates (6000 cells/well) and transfected as mentioned above. Transfected and non-transfected cells were incubated for 24, 48, and 72 h, respectively. Then cell proliferation was analyzed by MTT colorimetric assay. Experiments were performed thrice.

Cell apoptosis assay
After treatment, cells were washed, harvested, and counted. Then 1 × 10^5 cells were re-suspended in 100 μl binding buffer and incubated in the dark for 15 min at room temperature. Finally, 10 μl of Annexin V and 5 μl of PI (Sigma-Aldrich Co. LLC) were added according to the manufacturer’s instruction in the apoptosis kit (Biosea, Beijing, China). The apoptosis rate was determined with an Epics Altra II cytometer (Beckman Coulter, Danvers, USA). Cells were also viewed under an inverse fluorescent microscope. The experiment was repeated thrice.

Western blot analysis
Total protein was extracted and the concentrations were measured using a spectrophotometer (Bio-Rad, Hercules, USA). Then, sodium dodecyl sulphate–polyacrylamide gel electrophoresis was performed and transferred onto PVDF membrane (Millipore, Bedford, USA), followed by blocking with skimmed milk dissolved in tris buffer saline with tween-20 (TBST) for 1 h at room temperature. The membrane was incubated with primary antibody at 4°C overnight, and incubated with HRP-conjugated secondary antibody for 1 h at room temperature after washing thrice with TBST. After washing, the bands of protein were detected with ECL substrates (ZhongShan Co. Ltd, Beijing, China).

Cell cycle assay
After treatment, cells were harvested, washed with ice-cold PBS, and fixed with 70% ethanol at 4°C overnight. The ethanol was removed by centrifugation and ~10^6 cells were re-suspended in PBS containing 50 μg/ml PI and 50 μg/ml RNase A (Sigma-Aldrich Co. LLC) for 30 min in the dark before being analyzed on a FACScalibur flow cytometer (BD Biosciences, Bedford, USA). The percentage of cells at G0/G1, S, or G2/M phase was thereby calculated. DMSO-treated cells were used as control. Experiment was repeated thrice.

Quantitative real-time PCR
Cells were lysed with Trizol reagent (Invitrogen Tech) and total mRNA was extracted. The mRNA was reverse-transcribed into cDNA with a reverse-transcription kit (Promega Biotech). For PCR analysis, cDNA from triplicate dishes was diluted to a final concentration of 10 ng/μl. Quantitative real-time PCR was performed with a Universal Master Mix (Chembase, Beijing, China). cDNA (50 ng) was used to determine the relative amounts of mRNA by real-time PCR using MAX3000 Sequence-Detection System (Chambase) with specific primers and probes. The reaction was conducted for 40 cycles. β-actin was amplified as reference for stathmin. The primer and probe sequences are listed in Table 1.

Transwell assay
In the invasion assay, 50 μl Matrigel (BD Biosciences) was added to the upper chamber of the transwell apparatus with 8-μm pore size membrane (Costar, Cambridge, USA). After the Matrigel solidified at 37°C, cells were added into the upper chamber, and the lower chamber received completed medium. Membranes coated with Matrigel were swabbed with a cotton swab and fixed with 100% methanol for 10 min after 24 h incubation. The membranes with cells were soaked with crystal violet. The number of cells attached to the lower surface of the polycarbonate filter was counted. In the migration assay, all procedures were applied without Matrigel, and...
was performed in triplicate.

of the wound versus 0 h was calculated. Each experiment was monitored after 12 and 24 h, and the ratio of the width obtained from U87MG and U251 cell lines, respectively.

Two strains of GSCs, GSC-1 and GSC-2, were successfully obtained from U87MG and U251 cell lines, respectively (Fig. 1A). The stem cell phenotype of GSC-1 and GSC-2 was confirmed by western blot assay, which showed that the stem cell markers Oct3/4, MDR1, Sox2, and Nestin were up-regulated in the sphere-forming GSCs compared with the adherent cells (Fig. 1B).

**Table 1. Primers and probes used for qRT-PCR**

<table>
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<tr>
<th>Gene</th>
<th>Sequences</th>
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<tr>
<td><em>Stathmin</em></td>
<td>F: 5'-ACTGCCCTGTCGCTTGCTCT-3’&lt;br&gt;R: 5'-GTCCTGCAGCAGGGTCT-3’&lt;br&gt;P: 5'-CTTACAGTCGGTCGACGG-3’</td>
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<tr>
<td><em>β-Actin</em></td>
<td>F: 5'-CTCCATCTGGGCTCAGCTG-3’&lt;br&gt;R: 5'-GCTGTCACCTTCACCGTG-3’&lt;br&gt;P: 5'-CCAACACAGTGCTGTCTGG-3’</td>
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the cell number was calculated after 12 h. Experiments were done in triplicate.

**Scratch-wound healing recovery assay**

In wound healing assays, cell motility was assessed by measuring the movement of cells into a scraped area. The cells were placed in 24-well plates and scraped 2.0 mm wound with a 20 μl pipette tip. The speed of wound closure was monitored after 12 and 24 h, and the ratio of the width of the wound versus 0 h was calculated. Each experiment was performed in triplicate.

**VM assay**

Immediately before use, 24-well plates were coated with high-concentration Matrigel (200 μl/well) and incubated at 37°C for 40 min, until the Matrigel was solid. Cells were spun down, resuspended and seeded on Matrigel-coated wells at a density of 30,000 cells per well. human umbilical veins endothelial cells (HUVECs) were applied as reference. After incubation photomicrographs were taken for each well and the number of tubes (complete circular structures) was counted. The mean from the three readings of each well was used as the final reading.

**Statistical analysis**

Statistical analyses were performed using SPSS 16.0 statistical software (SPSS, Chicago, USA), and *P* < 0.05 was considered to show statistically significant difference.

**Results**

**Stathmin silencing suppressed proliferation and induced apoptosis of GSCs**

Two strains of GSCs, GSC-1 and GSC-2, were successfully obtained from U87MG and U251 cell lines, respectively (Fig. 1A). The stem cell phenotype of GSC-1 and GSC-2 was also confirmed by western blot assay, which showed that the stem cell markers Oct3/4, MDR1, Sox2, and Nestin were up-regulated in the sphere-forming GSCs compared with the adherent cells (Fig. 1B).

Application of siRNA to silence stathmin in GSCs resulted in reduced expression of stathmin at both mRNA and protein levels (Fig. 1C, *P* < 0.05). Cell proliferation was significantly suppressed 24 h after siRNA transfection (Fig. 1D, *P* < 0.05), whereas the rate of apoptosis was significantly increased (Fig. 1E, *P* < 0.05). Consistently, pro-apoptotic proteins (Bax and cleaved caspase-3) were up-regulated, while anti-apoptotic proteins (Bcl-xl and Bcl-2) were down-regulated in stathmin-siRNA transfected GSCs (Fig. 1F).

In the group of GSCs transfected with stathmin siRNA, the percentage of cells at G2/M phase was 36.4%, compared with 13.1% in the control group (Fig. 1G), indicating that the cell cycle was arrested at the G2/M phase with stathmin silencing.

Taken together, these results demonstrated that siRNA-mediated silencing of stathmin induces apoptosis and G2/M arrest in GSCs.

**Stathmin silencing suppressed the invasion and migratory abilities of GSCs**

We performed transwell analysis to examine whether the down-regulation of stathmin affects the invasion or migratory abilities of GSCs. The results showed that the invasive and migratory capacities of GSCs were significantly inhibited by stathmin silencing. The number of stathmin-siRNA transfected cells invading through the membrane was significantly lower than that of control-siRNA transfected cells (Fig. 2A,B, *P* < 0.05). Furthermore, cell migration was evaluated with a scratch-wound healing assay and the extent of cell migration into the scratched area was measured. The wounds in the wells of control-siRNA transfected cells healed rapidly and hardly a gap was left after 24 h. However, the wells of stathmin-siRNA transfected cells showed a much lower wound-healing ability than the control wells (Fig. 2C,D, *P* < 0.05). These data demonstrated that stathmin silencing suppresses both the invasion and migration capacities of GSCs.

**Stathmin silencing inhibited VM of GSCs**

The results of the VM assay showed both strains of GSCs had a strong ability of VM compared with the HUVEC group (Fig. 3A,B). However, the microtube formation capacity of GSCs was inhibited after stathmin-siRNA transfection (Fig. 3A); and GSCs which were transfected with siRNA formed fewer tubes compared with the control group (Fig. 3C, *P* < 0.05).

**Silencing of stathmin increased chemotheraphy sensitivity of GSCs to TMZ**

Transfection with stathmin siRNA significantly decreased the LD₅₀ for TMZ, an alkylating agent used for the treatment of GBM, from 1052.4 μM in control-siRNA transfected...
Stathmin silencing also significantly increased TMZ-induced apoptosis (Fig. 4B, \( P < 0.05 \)) and cell cycle arrest at G2/M phase (Fig. 4C) in GSCs. However, the apoptosis protein Bax and cleaved caspase-3 were up-regulated while anti-apoptosis protein Bcl-2 and Bcl-xl were down-regulated in the siRNA + TMZ group compared with the control group, TMZ group or siRNA group (Fig. 4D). These results demonstrated that silencing of stathmin increased chemotherapy sensitivity of GSCs to TMZ.

**Discussion**

Stathmin, known as metablastin, plays an important role in malignant cancers. The activity of stathmin is regulated by phosphorylation during the cellular transition from interphase to metaphase. The non-phosphorylated stathmin promotes the depolymerization of microtubules by sequestering tubulin [17] while the phosphorylated stathmin leads to increased microtubule stabilization and promotes the formation of mitotic spindles [18]. Stathmin is over-expressed in many malignant tumors, such as leukemia, non-small cell lung cancer, etc [19]. Stathmin over-expression can increase the invasion of prostate cancer, promote cancer progression and is associated with poor prognosis [20]. Silencing of stathmin can change the phenotype of malignant tumors, inhibit cancer cell proliferation and increase the chemotherapy sensitivity of cancer cells [21,22]. p53, which is a known tumor suppressor, can also modulate stathmin and induce cell cycle arrest at G2/M. Inhibition of stathmin in p53-mutant cell lines induces apoptosis [23]. In our previous work, we proved that stathmin expression in endothelial cells is associated with glioma WHO grade, and inhibition of stathmin suppresses proliferation,
invasion, and migration of vascular endothelial cells derived from glioma [16].

GSCs can self-renew and undergo multipotential differentiation, and are responsible for glioblastoma initiation, propagation, and recurrence [5]. In this study, we demonstrated that silencing of stathmin could inhibit proliferation, migration, invasion, and induce apoptosis of GSCs. GSCs are also the most important reason for resistance of chemotherapy. TMZ is a kind of nitrourea that could lead to DNA mismatches which could result in cell apoptosis in GBMs [24,25]. When
TMZ was applied, cell cycle was arrested at G2/M phase [26]. TMZ is very effective in clinical treatment, however, TMZ application induces modest increase of life span. The existence of GSCs is responsible for TMZ resistance [27]. As our result showed that the LD50 of TMZ in GSCs was significantly higher than that in adhesion cells which is around 100 μM [28]. Although application of siRNA targeting stathmin could enhance GSCs’ chemical therapy sensitivity to TMZ, but the LD50 was still as high as 800 μM. So how to eliminate GSCs is very important for glioma treatment.

In 1999, Maniotis et al. [10] were the first to report a new vascular entity named VM. The term VM has been used to describe the manner in which highly aggressive melanoma cells have the ability to form vasculogenic-like networks similar to embryonic vasculogenesis, which is independent of endothelial cells. The histological structures of VM channels are patterned networks of interconnected loops of periodic acid-Schiff (PAS)-positive extracellular matrix formed by aggressive tumor cells [29]. It is a new type vascularization of tumor that allows tumor cells to survive from hypoxia and denutrition environment. Moreover, a regular agent that targets VEGF for anti-angiogenesis therapies, such as endostar, works poorly for VM cure. In glioma, the VM-positive rate was associated with the grade of glioma, the GBM which is Grade IV has a strong ability of VM, and responsible for poor prognosis [9]. Thus developing a new way to inhibit VM may provide a potential method for GBM treatment. GSCs could also strongly induce endothelial cell migration and proliferation [30], as well as VM [14]. In this study, we demonstrated that silencing of stathmin could inhibit VM of GSCs. Taken together with our previous work [16], our data showed that inhibition of stathmin could suppress both angiopoiesis dependent on endothelial cells and VM dependent on aggressive tumor cells, which suggested that stathmin is a potential target for anti-angiopoiesis and anti-GSC treatment of glioma.

Further elucidation of the mechanisms of tumor angiogenesis and GSCs may provide more precise and effective anticancer therapies. Our findings suggested that stathmin plays an important role in glioma progression by supporting neo-angiogenesis. Stathmin may be used as a novel therapeutic target molecule in human glioma.

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