A simplified and modified procedure to culture brain glioma stem cells from clinical specimens

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Abstract. In recent years, the theory of brain glioma stem cells (BGSCs) has facilitated the study of gliomas. BGSCs have been accepted as the origin of gliomas and determine their biological features. Numerous efforts have been made to probe into the biological characteristics and behaviors of BGSCs. However, the culturing of target cells remains the essential first step for research on BGSCs. In this study, we established a simplified procedure to culture and isolate BGSCs from samples of clinical glioma patients. Samples of 17 glioma patients were included in the study, and the processed glioma cells were grown in serum-free stem cell media. After the tumor spheres appeared, a proliferation assay, a single-cell-derived colonies formation assay and an induced differentiation assay were carried out, followed by an immunocytochemistry assay. Serial passage was used to purify the target cells, whereas neither animal experiments nor sorting techniques were included. As a result, CD133\(^+\) BGSCs from 8 out of 17 patients were grown and maintained in a serum-free condition combined with EGF, FGF and B-27 supplements. The tumor sphere cells were serially passaged and showed pluripotency in an induced differentiation assay. Immunocytochemistry identified the committed markers (CD133, GFAP and TU-20) and confirmed the cells were BGSCs and their progeny. The results proved that CD133\(^+\) BGSCs from resected glioma tissue may be cultured in serum-free stem cell media, and may also be purified by conditioned culture combining serial passage, which is time-saving and cost-effective, and allows the cells to be used for subsequent research. The cell sorting techniques and animal experiments of tumorigenecity are optional. Thus, this modified procedure is more practical and feasible than other available procedures.

Introduction

As the most refractory tumors of the central nervous system, gliomas comprise 33.3-58.9% of intracranial tumors. The high-grade gliomas in particular (e.g., glioblastoma multiforme) have a dismal prognosis and a high recurrence rate; despite advances in multimodality therapy, the median survival is only 1 year (1,2). In recent years, the cancer stem cells theory (3) and the finding of glioma stem cells (4-6) have been the high-lights of glioma research. Brain glioma stem cells (BGSCs) have been proven to be the ‘seed’ cells of, and be responsible for, the initiation and development of gliomas, and thus have become new targets of glioma therapy (7). Currently the research on BGSCs mainly focuses on cell experiments, which demand a large number of cultured BGSCs as the first step. In this study, we introduce a simplified and modified procedure to culture, identify and purify BGSCs from resected samples of glioma patients, which is much more convenient and practical for the subsequent BGSC experiments.

Materials and methods

Tissue collection and grading. Brain glioma specimens were obtained within 30 min of surgical resection and were processed. The pathological grade of each specimen was confirmed by chief neuropathologists according to World Health Organization criteria. All 17 cases were inpatients in the Department of Neurosurgery of the First Hospital of China Medical University, China, between October 2008 and January 2009. The protocol of this study was approved by the Institutional Review Boards of the First Clinical Hospital, China Medical University.

Reagents and antibodies. Fetal bovine serum (FBS, qualifyed, Gibco, Invitrogen, Carlsbad, CA, USA), Dulbecco’s modified Eagle’s medium (DMEM)/F12 (Gibco), B-27 supplement (without serum and vitamin A, Invitrogen), recombinant human epidermal growth factor (rhEGF, Invitrogen), basic recombinant human fibroblast growth factor (rhFGF-b, Invitrogen), mouse monoclonal anti-CD133 antibodies (Abcam, Cambridge, UK), monoclonal rabbit anti-GFAP (glial fibrillary acidic protein) antibodies (Bioworld, Dublin, OH, USA), TU-20 (monoclonal mouse anti-β-tubulin III isoform, C-terminus, Millipore, Billerica, MA, USA), Cy3-conjugated goat anti-mouse secondary antibodies.
antibodies (Sigma, St. Louis, MA, USA), Cy3-conjugated goat anti-rabbit secondary antibodies (Sigma), fluorescein isothiocyanate (FITC)-conjugated goat anti-mouse antibodies (Sigma) and 4,6-diamidino-2-phenylindole (DAPI, Sigma) were used. The other common reagents were all analytically pure.

Culture of primary glioma cells and tumor spheres. Tumor-sphere cultures were performed according to reported procedures (4-6,8), with minor modifications. Following resection, the tumor tissues were washed, minced in phosphate-buffered saline (PBS) and subjected to enzymatic dissociation. The tissues were then mechanically dissociated with a graded series of fire-polished Pasteur pipettes, and passed through a series of cell strainers, and centrifuged at 800 x g for 5 min. Tumor cells were resuspended in DMEM/F12 containing 15% FBS and plated at a density of 2x10^5 live cells/ml. When tumor cells grew as monolayers in flasks, the medium was changed to committed stem cell medium (serum-free DMEM/F-12, 1:50 of B-27 supplement, 20 ng/ml rhEGF, 20 ng/ml rhFGF-b, 100 IU/ml penicillin G and 100 µg/ml streptomycin). Fresh rhEGF and rhFGF-b were added each week and the medium was changed twice a week. Cells were maintained in a standard tissue culture incubator with 5% CO_2 and 100% relative humidity at 37°C.

Proliferation assay and limited dilution assays. In the proliferation assay, when suspending cells emerged and the primary tumor spheres were clear, they were collected and dissociated into single-cell suspensions through a fire-polished Pasteur pipette, and then reseeded in the stem cell media at the same cell density. Each cell line was serially passaged over 4 generations. To evaluate the self-renewal capacity of tumor spheres, tumor spheres were mechanically dissociated into single-cell suspensions and reseeded into 96-well microwell plates by limited dilution at a cell density of 1-2 live cells per well. Each well was fed with fresh stem cell media every 3-4 days. The formation of cell clones was inspected under a phase-contrast microscope after 7-10 days.

Immunocytochemistry assays. The tumor spheres of approximately the 4th passage were collected and plated onto anti-peeling slides and incubated with stem cell media for 4 h. Following firm adhesion, the tumor spheres were fixed in 4% paraformaldehyde and incubated at 4°C with mouse monoclonal anti-CD133 antibodies diluted at 1:200 according to the manufacturer's instructions. Cy3-conjugated goat anti-mouse secondary antibodies diluted at 1:50 were added 24 h later and incubated for 2 h at room temperature.

To assess the multipotency of the tumor spheres, the tumor spheres were harvested and subjected to a differentiation assay by plating onto coverslips precoated with poly-L-lysine in DMEM/F-12 media containing 15% FBS and in the absence of growth factors or B-27 supplement. The media were changed every 2 days. The differentiated cells were fixed 7 days later, incubated with monoclonal rabbit anti-GFAP (1:200) for glial cells and TU-20 (1:200) for neurons overnight at 4°C. Secondary antibodies (Cy3-conjugated goat anti-rabbit and FITC-conjugated goat anti-mouse, 1:250) were then added and incubated for 2 h at room temperature.

For all of the staining, the cell nuclei were counterstained with DAPI for 5 min. In the control samples, primary antibodies were replaced by isotype IgG. A fluorescence microscope (Olympus BX61) was then used to observe and capture images of the results. The excitation wavelength was 554 nm for Cy3, 488 nm for FITC and 350 nm for DAPI.

Results

Tumor spheres culture. Tumor spheres were successfully cultured in 8 out of all 17 cases of glioma samples, including 2 primary glioblastoma multiformes (WHO grade IV), 1 recurrent anaplastic oligodendroglioma and 5 anaplastic astrocytomas (WHO grade III). The patients consisted of 3 males and 5 females. The ages varied from 44-73 years old, and the average age was 53.75 years old.

The monolayers formed 24-72 h after the single-cell suspensions were seeded in the serum-containing media. When switched into the stem cell media, single cell division occurred after 48-72 h, followed by the formation of large numbers of 'neurosphere-like' tumor spheres within 5-7 days. These tumor spheres were spherical or oval in suspending or semi-suspending states, with fine light refraction, consisting of 4-10 cells per sphere. Certain tumor spheres were of irregular morphology. The growth velocity was slow in the first few weeks, but within 2 weeks the majority of spheres had increased their diameters by 5-10-fold. Inspected under a microscope, the tumor spheres showed a smooth edge without prominent protrusion, and had poor light transmittance in the center as a result of a higher cell density. The majority of monolayer tumor cells still presented adherence, loss of proliferation, and subsequent differentiation, while tumor spheres remained suspended, continuing to proliferate and increase in cell numbers. Along with the growth of tumor spheres, a few adherent cells showed disaggregation, fragmentation and cell pyknosis, which manifested the progress of apoptosis. After approximately 2 weeks of culture, the formation of tumor spheres was observed and images were captured under a phase-contrast microscope (Fig. 1).

Proliferation and self-renewal assessment. In the proliferation assay, tumor spheres were dissociated into single-cell suspension and passaged at a ratio of 1:2 or 1:3. Cell cleavage occurred in 48 h and new tumor spheres formed within 1 week. Serial passage revealed the tumor spheres maintained a favorable proliferation ability after at least 4 generations. In a monoclonal formation assay, the single cells from tumor spheres were serially diluted and reseeded in microwells. Counted under a microscope, it demonstrated that more than 50% of the single cells in microwells were capable of forming new secondary tumor spheres, although the diameters were generally less than those of the primary spheres. Each new secondary sphere contained 10-40 cells, showing the same morphology as the parent sphere. This assay proved that individual cells from the parent tumor spheres were endowed with the ability to self-renew and form new secondary sphere colonies.

Immunocytochemistry identification. The tumor spheres were immunostained using CD133, the committed BGSC marker (4,8-11). After firmly attaching to the anti-peeling slides, an
immunocytochemical assay exhibited that the majority of tumor sphere cells were CD133-positive in a plasma membrane staining pattern under an immunofluorescence microscope. The nuclei were counterstained, exhibiting marked nuclear atypia. Images of the stained tumor spheres were captured, as shown in Fig. 2.

During induced differentiation, cell division in the spheres markedly decelerated and cells attached to the substrates, with cells migrating upon exposure to differentiation conditions. The tumor spheres became flat, exhibiting a radial morphology. The migrated cells formed monolayers, comprising various cell types with marked heteromorphism. Following differentiation in media with 15% FBS for 7 days, the cells were fixed and subjected to immunocytochemical detection. The results demonstrated that cells differentiated from the tumor spheres were positive for β-tubulin III and GFAP (Fig. 3), consistent with findings from other studies (4,10,11). The results indicated that CD133+ tumor spheres were multipotent for at least two neural cell types, neurons and astrocytes, which shows a multilineage differentiation ability.

Discussion

Gliomas are the most frequently diagnosed and aggressive primary intracranial tumors, with poor outcome. The mortality rate has remained on the increase in recent years; only less than 5% of patients survive the first 5 years, even after radical therapeutic strategies (12). Despite advances in glioma research and treatment in recent decades, prognosis has not substantially improved and the median survival period has been elevated by only 9-10 weeks (13). Thus, it is necessary and practical to find new targets and investigate new therapies based on glioma pathogenetic mechanisms. The theory of BGSCs provided new targets for the cure of gliomas (4-6,11,14). BGSCs are defined as a small population of cells capable of extensive proliferation, self-renewal, multipotent differentiation and tumor initiation. As the origins of glioma cells and the critical factors, BGSCs determine glioma propagation, progression and therapeutic resistance. Subsequently, BGSCs with a tumor-initiating ability have been isolated from GBMs, medulloblastomas, ependymomas and anaplastic oligoastrocytomas, and certain biological characteristics have been investigated (4,5,8,11,15,16). These findings suggest a hierarchical model in which glioma arises from BGSCs and progresses through mechanisms similar to a developmental process. It is well known that the eradication of gliomas can only be accomplished by targeting BGSCs. Nevertheless, little is known with regards to the biological mechanisms of BGSCs.

CD133, a 120 kDa cell-surface protein, which is a hallmark of normal human neural precursors, has been generally accepted as the marker of BGSCs regardless of whether they are solid tumors or glioma cell lines in vitro, although it is shared by other stem cells (4,9-11,16). In addition, some investigators used a flow cytometry-based side population (SP) technique to isolate BGSCs based on the characteristic that BGSCs were capable of excluding the fluorescent dye Hoechst 33342 (17). However, Hoechst 33342 is a type of liposoluble DNA-binding fluorescent dye and is cytotoxic to various cell types. Therefore, it is inappropriate to compare the respective biological features...
of sorted SP cells and non-SP cells. Thus, the SP sorting technique still calls for improvement. It is a key issue to find specific phenotype markers for BGSCs in subsequent studies, which may benefit further investigations on BGSCs.

The culture techniques for BGSCs are similar to the procedures for other stem cell types, such as serum-free media without adhesion factors and supplementation of rhEGF, rhFGF-β and B-27 (or N2 supplementation) (4-6,15,18,19). With regards to the sorting techniques, there are two basic methods available, one is immunomagnetic beads or fluorescent-activated cell sorting (FACS) based on the surface marker CD133; and the other method is the SP cell sorting technique based on a drug-resistant gene, such as an ATP-binding cassette or a multidrug resistance gene (MDR) (17,20). In either case, the supposed BGSCs are sorted, then identified and cultured. The procedures are complicated and expensive; following prolonged incubation, labeling, sorting and rinsing, the isolated BGSCs may not retain adequate viability for the subsequent culture process or relevant biological experiments. In this study, we simplified the steps of culture and identification to obtain expected BGSCs relatively rapidly and efficiently. The detailed culture process is the same as reported, but the isolation and identification are more convenient as described above. The results proved that the cultured tumor spheres could be serially passaged and possessed self-renewal ability, expressed the preferred BGSCs marker CD133, and differentiated into cells expressing β-tubulin III and GFAP, which were consistent with other findings (4,6,17). These results proved the ‘stemness’ of tumor spheres, therefore they could be defined as CD133+ BGSCs.

Compared with conventional methods, we did not use the sorting techniques (immunomagnetic beads, FACS, or SP techniques) or animal experiments to initiate new glioma. As is known, there are mainly 2 types of methods to purify BGSCs; conditioned purification or specific marker sorting combined with conditioned culture. Although target cells can be sorted using specific markers, the limitations remain clear, such as the long duration of the procedure, the high expense, complicated steps and loss of cell viability. Since the BGSCs were cultured in suspended tumor spheres, only the suspension cells were harvested and serially passaged during culture. Thus, the majority of non-BGSCs were excluded after they could be defined as CD133+ BGSCs.

A glioma growing environment that is extremely different to the microenvironment of spontaneous gliomas. Subsequently, the animal experiments were not always appropriate for the identification of BGSCs and were not included in our study, which saved time and expense.

Currently, BGSCs have attracted much attention and have been extensively investigated in fundamental fields such as BGSC biological characteristics, and clinical fields such as immunotherapy or gene therapy targeting BGSCs. Cytological experiments are substantial for BGSC research and obtaining enough viable BGSCs for experiments is a pivotal problem. Although numerous types of glioma cell lines were employed for BGSCs experiments, their biological features have significantly changed following long-term culture and serial passage in vitro and cannot mirror the primary characteristics of gliomas. Therefore, culturing BGSCs from clinical samples is an appropriate option. Nevertheless, the classical procedures comprise relatively complicated steps and a series of assays, and BGSCs finally cultured and isolated may not retain sufficient viability and natural properties as expected. In this study, we simplified the procedures and successfully grew BGSCs from resected glioma samples of clinical cases. The flow cytometry or immunomagnetic bead sorting methods were not used; conditioned culture and serial passage were used instead. The animal experiments for glioma tumorigenicity were also not performed due to controversies regarding their effectiveness. Within 3-5 generations, a considerable number of BGSCs were harvested with a high purity, which was confirmed by immunocytochemical identification of the tumor spheres. The results proved that this method is time-saving, cost-effective and convenient, contributing to the subsequent studies on BGSCs.

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References


