PC3-Secreted Microprotein Is Expressed in Glioblastoma Stem-Like Cells and Human Glioma Tissues

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Glioblastoma multiforme (GBM) is the most prevalent malignant primary brain tumor with a high recurrence rate. Despite multimodal therapy including surgical resection, chemotherapy, and radiotherapy, the median survival time after the initial diagnosis of GBM is approximately 14 months. Since cancer stem cells (CSCs) are considered the leading cause of cancer recurrence, glioblastoma stem cell-targeted therapy is a promising strategy for the treatment of GBM. However, because CSC heterogeneity has been implicated in the difficulties of CSC-target therapy, more in-depth knowledge of CSC biology is still required to develop novel therapies. In this study, we established single cell-derived tumorspheres from human glioblastoma U87MG cells. One of these tumorspheres, P4E8 clone, showed CSC-like phenotypes, such as self-renewal capacity, expression of CSC markers, resistance to anti-cancer agents, and in vivo tumorigenicity. Therefore, we used P4E8 cells as a cell-based model of glioblastoma stem cells (GSCs). Gene expression analysis using microarray indicated that the most highly expressed genes in P4E8 cells compared to the parental U87MG were PC3-secreted microprotein (MSMP). Furthermore, MSMP was expressed in patient-derived GSCs and human glioma tissues at the protein level, implying that MSMP might contribute to glioma development and progression.

Key words glioblastoma; cancer stem-like cell; PC3-secreted microprotein; tumorsphere; DNA microarray

INTRODUCTION

Glioblastoma multiforme (GBM) is the most common and aggressive primary brain tumor. The median overall survival of patients who received multimodal therapy including surgery, radiotherapy and chemotherapy, is approximately 14.6 months.1) Accumulating evidence indicated that cancer stem cells after surgery can cause a tumor recurrence.2) Therefore, strategies to isolate CSCs. This method was based on the resistance of stem cells to anoikis, a kind of apoptosis. When cell attachment or cell–matrix interaction was disrupted, anchorage-dependent differentiated cells induce anoikis, whereas stem cells survive and proliferate.10) However, anchorage-dependent cells can adhere together to survive and proliferate in floating cell culture conditions.11,12) Therefore, one promising approach to isolate CSCs is considered to form single-cell-derived tumorsphere because aggregation of anchorage-dependent cells must be prevented.13,14) In fact, this approach succeeded in the establishment of CSC lines from human colon cancer cell line,15) human primary colorectal cancer,16) human liver cancer,17) and human glioblastoma.18)

In this study, we established single cell-derived tumorspheres from human glioblastoma cell line U87MG, and one of the established tumorsphere, P4E8, showed CSC-like properties. To find novel therapeutic target molecules, we performed differential gene expression analysis between U87MG and P4E8 cells and identified that the most highly expressed instance, although CD133 was widely used as a CSC marker in glioblastoma, the CD133-negative cell population also exhibited stem cell characteristics and tumorigenicity.4,5) The SP assay is based on the high efflux ability of fluorescent dye in stem cells and several reports had indicated that stem-like cell population was isolated from cancer cell lines and tumor tissues by SP assay,6–8) but other report had shown that SP cells did not contribute to CSC phenotype.9)

On the other hand, sphere formation assay was also used to isolate CSCs. This method was based on the resistance of stem cells to anoikis, a kind of apoptosis. When cell attachment or cell–matrix interaction was disrupted, anchorage-dependent differentiated cells induce anoikis, whereas stem cells survive and proliferate.10) However, anchorage-dependent cells can adhere together to survive and proliferate in floating cell culture conditions.11,12) Therefore, one promising approach to isolate CSCs is considered to form single-cell-derived tumorsphere because aggregation of anchorage-dependent cells must be prevented.13,14) In fact, this approach succeeded in the establishment of CSC lines from human colon cancer cell line,15) human primary colorectal cancer,16) human liver cancer,17) and human glioblastoma.18)
gene in P4E8 cells was PC3-secreted microprotein (MSMP). Immunostaining analysis further indicated that the expression of MSMP in patient-derived glioma stem cells (GSCs) and human glioma tissues.

MATERIALS AND METHODS

Cell Culture Human glioblastoma U87MG cells were obtained from European Collection of Cell Cultures (ECACC, Wiltshire, U.K.) and were maintained in Minimum Essential Medium (MEM; Sigma, St. Louis, MO, U.S.A.) supplemented with 10% fetal bovine serum (FBS), 1% non-essential amino acid (Gibco, Thermo Fisher Scientific, Waltham, MA, U.S.A.), 1 mM sodium pyruvate (Nacalai Tesque, Kyoto, Japan), and 100 U/mL penicillin, 100 µg/mL streptomycin (Invitrogen, Thermo Fisher Scientific) in humidified air atmosphere containing 5% CO2 at 37°C. Single cell-derived U87MG clones and previously established patient-derived GSC lines (146, MDI3) (kind gifts from Dr. Ichiro Nakano, University of Alabama at Birmingham) were cultured as floating spheres in neurosphere medium (Dulbecco’s modified Eagle’s medium/F12 supplemented with 1% N2 Max Media Supplement (R&D Systems, Minneapolis, MN, U.S.A.), 2% B27 Supplement Minus Vitamin A (Thermo Fisher Scientific), 3 mM sodium bicarbonate, 100 U/mL penicillin, 100 µg/mL streptomycin, 20 ng/mL basic fibroblast growth factor and epidermal growth factor (EGF). Human embryonic stem cell (hESC) (Thermo Fisher Scientific) were cultured according to the manufacturer’s protocol.

Establishment of Single Cell-Derived Tumorspheres from U87MG Cells U87MG cells dissociated with Accutase (Innovative Cell Technologies, San Diego, CA, U.S.A.) were washed twice with ice-cold 2% FBS-phosphate-buffered saline (PBS) and stained with 7-aminoactinomycin D (7-AAD) (BD Biosciences, Franklin Lakes, NJ, U.S.A.) to exclude dead cells. Cells were first gated by forward side scatter (FSC-A vs. SSC-A) to exclude cellular debris. Single cells were then separated from doublets by two subsequent gates in plot of FSC-A vs. FSC-W and SSC-A vs. SSC-W. Finally, 7-AAD-negative live cells were sorted into 150 µL of neurosphere medium in 96-well suspension culture plates (Sumilon, Sumitomo Bakelite, Tokyo, Japan) using a FACS Aria II (BD Biosciences, Franklin Lakes, NJ, U.S.A.) to exclude dead cells. Cells were first gated by forward side scatter (FSC-A vs. SSC-A) to exclude dead cells. Single cells were then separated from doublets by two subsequent gates in plot of FSC-A vs. FSC-W and SSC-A vs. SSC-W. Finally, 7-AAD-negative live cells were sorted into 150 µL of neurosphere medium in 96-well suspension culture plates (Sumilon, Sumitomo Bakelite, Tokyo, Japan) using a FACS Aria II (BD Biosciences, Franklin Lakes, NJ, U.S.A.). After two weeks, the phase contrast image of each sphere was observed, and the diameters were measured using a Leica AF7000 fluorescence microscope (Leica, Heerbrugg, Switzerland). Individual spheres were expanded and used for further experiments.

Xenotransplantation All animal experiments were approved by the Animal Care and Use Committee of Kansai Medical University (No. 14-121) and performed in accordance with the institutional guidelines. Six- to eight-week-old female BALB/c nude mice (BALB/c slc-nu/nu, Shimizu Laboratory Supplies Co., Ltd., Kyoto, Japan) were used in this study. After U87MG cells and tumorspheres were dissociated with Accutase, cells (3 × 105 or 1 × 106) in 100 µL of PBS were subcutaneously injected into the left flanks of the mice. The same amount of PBS was injected into the right flank as a control. Tumor volume was measured every 5 d and calculated using the following formula: \[ V = \frac{1}{2} D \times d^2 \] where D and d are the longest and the shortest diameters, respectively. To assess tumor-initiating ability, mice bearing tumors larger than 100 mm3 were counted as tumor-bearing mice.

For the orthotopic implantation, cells (5 × 105 cells in 2 µL of PBS) were stereotactically injected into the right striatum of mice under anesthesia with the mixture of midazolam, medetomidine, and butorphanol. After 14 d, the mice were deeply anesthetized and sacrificed with carbon dioxide. The mice were then perfused transcardially with PBS, followed by 4% paraformaldehyde. The brain was dissected out and immersed in 4% paraformaldehyde overnight at 4°C. After embedded in paraffin, coronal sections (5 µm in thickness) were prepared and stained with hematoxylin–eosin (H&E) staining to indicate the tumor area.

Immunocytochemistry Cells seeded onto 8-well chamber slides (Matsunami Glass Industry, Osaka, Japan) (double-coated with 15 µg/mL poly-1-ornithine and 10 µg/mL laminin) were fixed with 4% paraformaldehyde and permeabilized with 0.2% Triton X-100. After blocking with Blockace (Dainippon Pharmaceutical, Osaka, Japan), cells were incubated with primary antibodies at 4°C overnight, followed by the appropriate secondary antibodies. Antibodies used are listed in Supplementary Table 1. Cell nuclei were stained by incubation with 2 µM Hoechst33342 for 30 min at room temperature, following treatment with secondary antibody. Fluorescence images were acquired by the LSM 700 confocal microscope (Carl Zeiss Microscopy GmbH, Jena, Germany).

Limiting Dilution Assay Limiting dilution assay was performed as previously described. Briefly, cells were dissociated with Accutase and plated on 96 well plates in two-fold serial dilution ranging from 0.98–500 cells/well in neurosphere medium. After 6d, the number of wells without spheres were counted and plotted against the number of cells per well. The number of cells required to form at least one tumorsphere was calculated from the point at which 37% of the well did not have tumorspheres.

Cell Viability Assay Cells were seeded onto poly-1-ornithine/laminin (Sigma-Aldrich, St. Louis, MO, U.S.A.)-coated 96-well plates at a density of 3000 cells per well. After 24 h, the medium was changed to a growth medium containing 0.2% dimethyl sulfoxide (DMSO) and 100 µM of temozolomide, cisplatin, etoposide, nimustine, 5-fluorouracil, methotrexate, tamoxifen, all-trans retinoic acid (Wako Pure Chemical Corporation, Osaka, Japan) and irinotecan (LKT Laboratories, St. Paul, MN, U.S.A.). After 3 d, 10 µL of Cell Counting Kit-8 (Dojindo, Kumamoto, Japan) reagent was added to each well, and cells were incubated for 4 h at 37°C. The absorbance was then measured at 450 nm using a 2030 ARVO X Multilabel Reader (PerkinElmer, Inc., Life and Analytical Sciences, Waltham, MA, U.S.A.).

RT-PCR and Real-Time Quantitative PCR (qPCR) Total RNA was extracted with TRIzol reagent (Ambion, ThermoFisher Scientific) and purified using the PureLink RNA Mini Kit (Ambion, ThermoFisher Scientific). First-strand cDNA was synthesized from 2 µg of total RNA using SuperScript III reverse transcriptase (Invitrogen, Thermo Fisher Scientific) with oligo(dT)12-18 primer (Invitrogen) in a volume of 20 µL and used as template cDNA for subsequent PCR reaction.

RT-PCR was performed with template cDNA (30 ng), ExTaqHS (TaKaRa Bio, Shiga, Japan) in a 25 µL reaction using a Mastercycler Gradient thermal cycler (Eppendorf, Hamburg, Germany) according to the following protocol: ini-
tial denaturation at 94 °C for 30 s, followed by 20–34 cycles of denaturation at 98 °C for 10 s, annealing at 58–60 °C for 30 s and extension at 72 °C for 45 s. The final extension was carried out at 72 °C for 3 min. Specific primers (Fasmac, Kanagawa, Japan), annealing temperature, and the number of cycles for each reaction are listed in Supplementary Table 2. The absence of genomic contamination was confirmed by using reverse transcriptase-negative samples.

Real-time qPCR was carried out using StepOne Real-Time PCR system (Applied Biosystems, Foster City, CA, U.S.A.). The PCR reaction mixture consists of first-strand cDNA (20 ng), 10 μL of FastStart Universal Probe Master (Roche Applied Science, Indianapolis, IN, U.S.A.), 200 nM gene-specific forward and reverse primers (Fasmac), and 200 nM Universal Probe Library Probes (Universal Probe Library Human Set, Roche Applied Science) in 20 μL reaction volume. PCR conditions were initially set at 95 °C for 10 min, followed by 50 cycles of 95 °C for 15 s and 60 °C for 1 min. Specific primers and Universal Probes were listed in Supplementary Table 3. Experiments were performed in triplicate using the delta–delta Ct method, and glyceraldehyde-3-phosphate dehydrogenase was used as an endogenous control to normalize expression data.

**DNA Microarray** The integrity of total RNA isolated from U87MG (n = 3) and P4E8 (n = 3) cells was assessed using a BioAnalyzer 2100 (Agilent Technologies, Waldbronn, Germany), and no degradation was observed. Microarray analysis was performed by Hokkaido System Science (Sapporo, Japan) using the SurePrint G3 Human 8x60K ver. 2.0 (Agilent, Santa Clara, CA, U.S.A.). The data analysis was performed with GeneSpring GX. The raw data were normalized with a 75 percentile shift, and baseline transformation was performed using the median of all samples. Subsequently, the relative expression of each gene was analyzed. The data have been deposited in the NCBI Gene Expression Omnibus (GEO) database (GSE146698).

**Immunohistochemistry** This study was performed following the ethical principles for medical research outlined in the Declaration of Helsinki and relevant guidelines and regulations. Written informed consent was obtained from each patient and this study was approved by the Institutional Review Board of Kansai Medical University (No. 1401-1, 2017085).

Tumor specimens were obtained from patients with glioma by surgical resection at the department of neurosurgery, Kansai Medical University. Formalin-fixed paraffin-embedded blocks of human glioma tissues were cut into 5 μm-thick sections and mounted on glass slides. Sections were deparaffinized in xylene and rehydrated in a graded series of ethanol. The sections were then immersed in 20 mM sodium citrate buffer (pH 6) and antigen retrieval was performed by autoclave heating at 120 °C for 20 min. After blocking with Blockace, sections were incubated overnight at 4 °C with rabbit polyclonal antibodies against Sox2 (1:1000 dilution) or MSMP (1:200 dilution) (Supplementary Table 1), followed by treatment with anti-rabbit immunoglobulin G (IgG) biotinylated goat IgG (1:100 dilution, Vector Laboratories, Burlingame, CA, U.S.A.). Then, sections were incubated with avidin-biotin-peroxidase complex (ABC) reagent (VECTASTAIN ABC-Kit, Vector Laboratories) and visualized using ImmPACT DAB (Vector Laboratories). Images were captured using a Nikon Eclipse Ni light microscope (Nikon, Tokyo, Japan) under bright-field illumination.

**Statistical Analysis** The data for tumor growth in vivo were analyzed by two-way repeated measures ANOVA with Bonferroni post hoc test using GraphPad Prism v.4.03 software (GraphPad Software Inc., San Diego, CA, U.S.A.). The comparison of drug sensitivity between U87MG and P4E8 cells was performed by two-sided Welch’s t-test using Microsoft Excel version 2013.

**RESULTS**

**Establishment of Single Cell-Derived Tumorsphere from U87MG Cell Line** It had been reported that CSC-like cells were isolated from several human glioblastoma cell lines including U87MG cells, which was widely used in brain tumor research.21–23 To isolate CSC-like population from U87MG cells, the expression of CSC marker CD133 was first examined by a flow cytometer because CD133 expression was controversial in U87MG cells.24–26 As shown in Supplementary Fig. 1, the CD133 expression in U87MG cells was not observed. Then, we examined the existence of side population (SP) cells by FACS analysis under adherent and sphere culture condition (Supplementary Fig. 2) because sphere culture was widely used to enrich CSCs. However, since verapamil, an inhibitor of Hoechst33342 transport, did not significantly change the SP cell population in adherent and sphere culture of U87MG cells, it is concluded that CD133-positive cells and SP cells...
were not present in U87MG cells.

Therefore, a single cell-derived sphere formation was employed to isolate the CSC-like population from U87MG cells. Single cells were sorted into 96-well plates (1 cell/well) by FACS and were further incubated in a neurosphere medium for 2 weeks (Fig. 1A). As shown in Fig. 1B, the sphere formation efficiency for \( >100 \) and \( >50 \mu m \) tumorsphere diameter were 3.3 ± 0.3 and 9.2 ± 1.3 %, respectively. Spheres for \( >100 \mu m \) tumorsphere diameter were further expanded separately, and 14 of single cell-derived tumorspheres were established.

To characterize the established tumorspheres, gene expression of known CSC markers was analyzed by RT-PCR. As shown in Fig. 2A, the marker expression profile was varied for each tumorsphere. All markers examined were expressed in 6B4, P4E8, P5F4, P5G5, and tumorspheres derived from the bulk U87MG cells (bulk tumorsphere). Among these tumorspheres, especially high expression of both Sox2 and Nestin were seen in P4E8 and P5G5 tumorsphere. These tumorspheres also had a shorter doubling time (31.7 ± 0.9 h for P4E8, 34.8 ± 0.3 h for P5G5) than others (Fig. 2B). The mRNA expression of Sox2 and Nestin in bulk tumorsphere was slightly higher than that in adherent U87MG cells but lower than in P4E8 and P5G5. These results suggest that bulk tumorsphere was composed of sphere forming cells with distinct CSC marker expression profiles, but the floating culture of the single cell could establish clonal tumorsphere.

**Tumorigenic Ability of Established Single Cell-Derived Tumorspheres** To evaluate the in vivo tumorigenicity, P4E8 and P5G5 tumorspheres were subcutaneously injected into the nude mice because these tumorspheres highly expressed Sox2, which had been reported to be important for tumorigenicity and proliferation of glioblastoma initiating cells. \(^{27} \) For comparison, U87MG cells under adherent culture condition were also investigated. When \( 1 \times 10^6 \) cells were injected, P4E8 cells generated significantly larger tumors than U87MG cells on days 25, 30, and 35 (Figs. 3A, B). Even inoculation of \( 3 \times 10^5 \) P4E8 cells tends to form larger tumors than \( 1 \times 10^6 \) U87MG cells. On the other hand, P5G5 cells did not generate a tumor until day 35. Furthermore, orthotopic injection of U87MG and P4E8 cells into the brains of nude mice was carried out (Fig. 3C). At 14d after injection, P4E8 cells formed brain tumors in three out of four mice, but adherent U87MG cells did not.

These results indicated that P4E8 cells have highly tumorigenic and proliferative properties compared to adherent U87MG cells.

**Characterization of CSC-Like Properties of P4E8 Tumorsphere** To further characterize the CSC-like properties...
Tumor incidence is displayed in the upper right corner of each image. (Color)

Representative images of H&E-stained coronal brain sections at 14 d after intracranial injection with U87MG or P4E8 cells (*mice inoculated with P4E8 (1 × 10^6) cells compared to U87MG cells (1 × 10^5) (p < 0.001). (B) Representative pictures of mice at 35 d after injection with indicated number of U87MG or P4E8 cells were shown. Arrowheads indicate tumors.}

Adherent U87MG, P4E8 and P5G5 cells were subcutaneously injected into the left flank of Balb/C nude mice at indicated cell number (n = 4 mice in each group). (A) After injection, each tumor volume was measured every 5 d. Data represents the mean ± S.D. from four mice per group. Two-way repeated measures ANOVA with Bonferroni post hoc test showed significant difference of tumor growth of at least one tumorsphere/well was calculated as 29.7 cells in U87MG cells and 2.6 cells in P4E8 cells, indicating that P4E8 tumour had higher self-renewal ability than U87MG cells (Fig. 4B). Furthermore, P4E8 cells increased resistance against anti-cancer agents, cisplatin, etoposide, 5-fluorouracil, methotrexate, tamoxifen, and all-trans retinoic acid, compared with U87MG cells (Fig. 4C), but only temozolomide enhanced cytotoxicity on P4E8 cells. Among these agents, tamoxifen, an anti-estrogen agent, showed the highest cytotoxic activity against P4E8 cells.

Taken together, these results indicated that P4E8 cells have CSC-like properties such as expression of CSC markers, self-renewal ability, and resistance against various anti-cancer agents, in addition to the high tumorigenic ability. Therefore, P4E8 cells were used as a cell-based model of GSCs for further experiments.

**Differential Gene Expression Analysis between Parental U87MG Cells and P4E8 Tumorsphere** To find new therapeutic targets for CSCs, differentially expressed genes between adherent U87MG and P4E8 tumorsphere were analyzed by cDNA microarray. A total of 3081 genes were differentially expressed at a two-fold cut-off with p-value <0.05, and 1309 upregulated and 1686 downregulated genes were identified (Fig. 5A). The top 10 significantly upregulated genes in P4E8 cells were shown in Table 1. Among these genes, the highest expressed gene was **MSMP**, and the second highest expressed gene was solute carrier family 24 member 3 (**SLC24A3**). A known CSC marker, Sox2, was also upregulated in P4E8 cells, supporting the validity of our data. On the other hand, the high expression of the T cell receptor beta constant 1 (**TRBC1**) gene in P4E8 cells would be false positive or artifact because TRBC1 is a T cell receptor component in T lymphocyte. To validate the microarray data, real-time qPCR was performed for **MSMP**, **SLC24A3**, Phospholipase A2 receptor 1 (**PLA2R1**), FYVE, RhoGEF and PH domain containing 4 (**FGD4**), Prickle homolog 1 (**PRICKLE1**), Sox2, Contactin-associated protein-like 3 (**CNTNAP3**), and Potassium voltage-gated channel interacting protein 3 (**KCNIP3**) (Fig. 5B). It is difficult to distinguish between **CNTNAP3** and **CNTNAP3B** gene by PCR analysis because these genes are paralogs with high sequence similarity; therefore, **CNTNAP3B** transcripts would also be amplified by the primer set for **CNTNAP3**. As shown in Fig. 5B, all transcripts were highly expressed in P4E8 cells compared with U87MG cells, supporting the microarray results, although **SLC24A3**, **PRICKLE1**, and **KCNIP3** transcripts were undetected in U87MG cells. In addition, when the expression of these transcripts in P4E8 cells was compared with that in human neural stem cells (hNSC), used as a normal stem cells, the expression of **MSMP**, **SLC24A3**, **PLA2R1**, and **KCNIP3** transcripts were higher in P4E8 cells than in hNSC.

**Expression of MSMP in Patient-Derived GSC Lines and Human Glioma Tissues** Recent evidence showed that GBM could be classified into at least four molecular subtypes. Among these subtypes, proneural and mesenchymal subtypes were mutually exclusive and the mesenchymal subtype displayed a more malignant phenotype than proneural subtype. In the previous report, patient-derived GSCs, 146 and MD13, were molecularly characterized and assigned to proneural and mesenchymal subtype, respectively.

First, the CSC marker expression pattern was compared of P4E8 cells, the expression of known CSC markers (Sex determining region Y (SRY)-Box2 (Sox2), stage-specific embryonic antigen 1 (SSEA-1), and Nestin) and differentiation markers (S100β, glial fibrillary acidic protein (GFAP), and βIII tubulin) was examined by immunocytochemistry. As shown in Fig. 4A, higher expression of Sox2 and SSEA-1 was observed in P4E8 cells than in U87MG cells. On the other hand, Nestin was expressed in both U87MG and P4E8 cells. For astrocytic markers, S100β was expressed in both U87MG and P4E8 cells, but GFAP was not. For neuronal marker βIII tubulin, weak signals were partially seen in both U87MG and P4E8 cells. Next, the sphere-forming frequency was determined by limiting dilution assay. The number of cells required to generate
between P4E8 cells and patient-derived GSCs by immunocytochemistry. SSEA1, highly expressed in P4E8 cells (Fig. 4A), was not expressed in both MD13 and 146 cells (Fig. 6A). On the other hand, Sox2 was more highly expressed in 146 cells than in P4E8 and MD13 cells (Figs. 4A, 6A). These results indicated that the CSC marker expression pattern was different among P4E8, proneural and mesenchymal GSCs.

Next, we determined the expression of MSMP, SLC24A3, PLA2R1, and KCNIP3 in patient-derived GSCs. Among these genes, mRNA expression of MSMP, SLC24A3 and KCNIP3 were detected in 146 and MD13 cells, but PLA2R1 transcripts was not (Supplementary Fig. 3A). We further examined the protein expression of SLC24A3 and KCNIP3 by Western blot analysis. As shown in Supplementary Fig. 3B, a significant expression of SLC24A3 was detected in MD13 cells compared with that in P4E8 cells. For KCNIP3, the expression was not detected in the examined cell lines (Supplementary Fig. 3C).

The protein expression of MSMP was examined in U87MG, P4E8, and patient-derived GSCs. As shown in Fig. 6B, MSMP showed a punctate pattern in the cytosol of all these cells examined, indicating the localization of MSMP in secretory vesicles (Fig. 6B). Although strong signals were observed in P4E8 and MD13 cells, MSMP was also expressed in U87MG cells. These results suggested that MSMP would be expressed in both GSC and non-GSC but at higher levels in GSC.

Finally, MSMP expression was examined in human glioma tissues graded from I to IV by immunohistochemistry. As shown in Fig. 7, MSMP was detected in the cytoplasm in a punctate pattern in all grade glioma. On the other hand, the expression of Sox2 was higher in high-grade glioma (grades III, IV) compared with low-grade glioma (grades I, II). These results indicated that MSMP was actually expressed in human glioma tissues regardless of glioma grade.

Fig. 4. Characterization of CSC-Like Properties in P4E8 Cells

(A) Immunocytochemical analysis for marker expression in U87MG and P4E8 cells. The fluorescence signals derived from CSC markers (Sox2, Nestin, SSEA-1), astrocyte markers (S100β and GFAP) and neuronal marker (βIII tubulin) were indicated by green. Nuclei were stained with Hoechst33342 (red pseudo-color). Arrowheads indicate Sox2 positive cells. Scale bars, 50µm. (B) In vitro limiting dilution assay of U87MG and P4E8 cells. (C) Drug sensitivity against various chemotherapeutic drugs in U87MG and P4E8 cells. Cells were treated with indicated anti-cancer agents at 100 µM for 3 d. Data represent the mean ± S.D. of three separate experiments. Statistical difference was determined by Welch’s t-test. Asterisks indicate significant difference between U87MG and P4E8 cells (*p < 0.01). (Color figure can be accessed in the online version.)
Fig. 5. Differential Gene Expression Analysis between U87MG Cells and P4E8 Tumorsphere by DNA Microarray and Validation of Microarray Data by Real-Time qPCR

(A) The heat map shows 1395 upregulated (>2-fold) and 1686 downregulated (<2-fold) genes in P4E8 tumorspheres compared to U87MG cells in biological triplicates. (B) Validation of DNA microarray experiments by real-time qPCR. In addition to U87MG and P4E8 cells, hNSC was used as a normal stem cells. Each bar indicates fold-increase of transcripts compared with the level of expression in U87MG or hNSC (light gray), which is set at a value of 1. Data represent the mean ± S.D. from three separate experiments. ND, not detected. (Color figure can be accessed in the online version.)

Table 1. Top 10 Genes Up-Regulated in P4E8 Cells (versus U87MG Cells)

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<th>Accession No.</th>
<th>Descriptions</th>
<th>Fold change</th>
<th>p-Value</th>
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<td>NM_001044264</td>
<td>Microseminoprotein, prostate associated (MSMP)</td>
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<td>SRY-box 2 (SOX2)</td>
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<tr>
<td>NM_139241</td>
<td>FYVE, RhoGEF and PH domain containing 4 (FGD4)</td>
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<td>&lt;0.0001</td>
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<td>NM_013434</td>
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<td>Phospholipase A2 receptor 1, 180kDa (PLA2R1)</td>
<td>112.7</td>
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DISCUSSION

Targeted therapies against CSCs are considered to be a promising therapeutic strategy. However, identification and isolation of CSCs from tumor tissues have still been challenging because of a rare population and phenotypic heterogeneity of CSCs. Even in the same human GBM tissue, CSCs derived from the periphery or core region had distinct properties such as proliferation potential and tumor-initiating ability.\(^{29,30}\) Since CSC heterogeneity is also implicated in the difficulties of CSC-targeted therapy,\(^{31}\) exploring more target molecules of CSCs is important for further clinical implication.

We established tumorspheres from human glioblastoma U87MG cells by the floating culture of single cell in the neurosphere medium. Among established tumorspheres, P4E8 cells were considered a cell-based model of GSCs because P4E8 cells showed CSC-like phenotypes such as self-renewal ability, expression of CSC markers, drug resistance, and high tumorigenic potential.

CSCs were known to be resistant to anti-cancer drugs. One of the cause of drug resistance is an overexpression of ATP-binding cassette (ABC)-transporter family such as P-glycoprotein (P-gp), multidrug resistance-associated proteins (MRPs) and breast cancer resistance protein (BCRP).\(^{32,33}\) We showed that P4E8 cells were resistance to cisplatin, etoposide, 5-fluorouracil, methotrexate, tamoxifen and all-trans retinoic acid (Fig. 4C). Among these agents, cisplatin, etoposide, 5-FU, methotrexate, tamoxifen had been reported to be substrates of ABC transporters.\(^ {34}\) Although temozolomide and SN-38 (the active metabolite of irinotecan) had been reported to be substrates of ABC transporer,\(^ {34,35}\) P4E8 cells did not show the drug resistance against these agents. These results suggested the contribution of particular ABC transporters to drug resistance in P4E8 cells. On the other hand, since significant cytotoxic activity against P4E8 cells was observed in treatment with tamoxifen (Fig. 4C), this drug might be a good candidate for GSC-targeted therapy. Thus, the cell-based model of CSC is useful for screening for CSC-targeting agents. In addition, single cell-derived tumorsphere formation would also be useful for isolation of CSC from tumor tissues by not being...
dependent on the expression of cell surface markers.

As shown in Fig. 2A, both P4E8 and P5G5 showed higher mRNA expression of Sox2 compared with other established tumorspheres, but only P4E8 cells formed tumors (Fig. 3A). It had been reported that the Sox2 protein function were regulated through various post-translational modifications such as phosphorylation, acetylation, methylation, SUMOylation, and ubiquitylation. Therefore, the different post-translational modifications of Sox2 between P4E8 and P5G5 cells may contribute to the different tumorigenicity in vivo. Sox2 is known to be a CSC marker of proneural subtype but not of mesenchymal subtype. Immunochemical analysis revealed that the expression of Sox2 in P4E8 cells was comparable to mesenchymal MD13 cells but was lower than that in proneural 146 cells (Figs. 4A, 6A). These results suggest that P4E8 cells would not be a proneural subtype.

We identified highly expressed genes in P4E8 cells compared to the parental U87MG by microarray analysis. Among these genes, the expression of MSMP, SLC24A3, PLA2R1, and KCNIP3 transcripts were higher in P4E8 cells than in normal hNSC, but the expression of PLA2R1 transcripts and KCNIP3 protein were undetected in patient-derived GSCs (Supplementary Fig. 3). On the other hand, a significant expression of SLC24A3 was detected in MD13 cells compared with that in P4E8 cells. Although further studies are required, these results suggest that SLC24A3 may be involved in the function of mesenchymal GSCs.

The human MSMP gene, encoding an 11.1 kDa secretory protein, had been identified by sequence similarity to human β-microsemionprotein. Although the expression of MSMP mRNA in normal human tissues was limited to trachea and testis, MSMP transcripts and protein were highly expressed in prostate cancer cell line PC-3. In addition, the immunohistochemical analysis had indicated the high level expression of MSMP in human benign prostate hyperplasia, prostate cancers and breast tumor tissues. This study first indicated that MSMP was expressed in patient-derived GSCs (146 and MD13) and human glioma tissues.

The secretion of endogenous MSMP had previously been determined in human prostate cancer PC-3 cells and human ovarian cancer cell lines. To determine the secretion of MSMP from U87MG, P4E8, and PC-3 cells, culture medium was harvested after 48 h incubation of 1 × 10^6 cells in 6 well plates and the enzyme-linked immunosorbent assay was performed (data not shown). As a result, the secreted MSMP was undetectable level in the culture medium of all these cell lines including PC-3 cells. Since MSMP has been well known to be secreted from PC-3 cells, further detailed investigation of experimental condition is required to determine the secretion of MSMP in U87MG and P4E8 cells. On the other hand, it had been reported that MSMP expression and secretion was low under normoxic condition but high under hypoxic condition in ovarian cancer cell lines, suggesting that the secretion of MSMP from P4E8 cells may also be induced under hypoxic condition.

Although the functions of MSMP are not fully understood, it had been reported that MSMP had a chemotactic activity toward peripheral blood monocytes through C-C chemokine receptor 2 (CCR2), the receptor of monocyte chemotactic protein 1 (MCP-1). In solid tumors, tumor-associated macrophages (TAMs), a component of the tumor microenvironment, have been implicated in promoting tumor progression and associated with poor prognosis. Therefore, it is speculated that MSMP expressed from glioma cells, including GSCs, may increase the recruitment of monocytes from the circulation, resulting in the increase of the number of TAMs and enhancement of glioma progression.

Although the functional analysis was required to elucidate the role of MSMP in glioma progression, our findings raise the possibility that MSMP may contribute to glioma development or progression, or both.

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Conflict of Interest The authors declare no conflict of interest.

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