Extended Neurosphere Culture of Brain Tumor Stem Cells with the PromoCell Cancer Stem Cell Medium

Application Note

The PromoCell Cancer Stem Cell Medium

While adherent cultures of brain tumor cells in fetal calf serum-containing standard media undergo changes in the form of partial differentiation, 3D cultures of brain cancer cells as neurospheres in serum-free conditions maintain – and therefore reflect better – the properties of primary tumors [1]. However, conventional serum-free media for 3D culturing of neural cancer stem cells as neurospheres struggle to efficiently maintain self-renewing driver subpopulations of cancer stem cells (CSCs). Consequently, the stem cell population in the culture is gradually lost along with its proliferation potential (see Fig. 1, grey chart).

The PromoCell Cancer Stem Cell Medium is designed to meet your requirements for undifferentiated and extended serial 3D neurosphere culture of most commonly used brain tumor cell lines. Its advanced and defined formulation permits 3D expansion as self-renewing neurospheres while maintaining rigorous cellular selection for stem cell traits, i.e. self-renewal and anoikis resistance. This results in an optimal culture environment for brain CSCs, as is reflected in the sustained serial passage ability of 3D neurospheres with the Cancer Stem Cell Medium. Thus, efficient neurosphere expansion allows to easily access cells enriched for sharing these aforementioned stem cell properties (see Fig 1, red chart).

PromoCell’s Cancer Stem Cell Medium is defined and ready to use, providing a standardized culture devoid of stimuli of uncharacterized origin. This is a significant benefit in terms of neural stem cells, which are highly responsive stem cells that require a reliable and reproducible control of the self-renewal/differentiation axis. The Cancer Stem Cell Medium is suitable for a cost-effective, standardized routine culturing of neural cancer cell lines as neurospheres. In contrast to the limited numbers of passages achieved with conventional serum-free formulations, the Cancer Stem Cell Medium supports long-term passageable neurosphere cultures of a broad variety of cell lines.

Fig. 1: Plot of cumulative population doublings of U-87 MG glioblastoma cells during cultivation as 3D neurospheres in the Cancer Stem Cell Medium compared to an established commercial neurosphere culture medium. The Cancer Stem Cell Medium maintained neurosphere formation and steady proliferation during the entire culture, which was discontinued after passage 11 with no signs of growth rate inhibition. By comparison, when cultured in the competitor neurosphere culture medium, a steadily declining expansion rate of U-87 MG was observed; this was indicative of the progressive loss of the primitive self-renewing driver cell population.
Background

The term “brain cancers” encompasses a variety of malignancies of the central nervous system (CNS), e.g. gliomas, meningiomas, pituitary adenomas and nerve sheath tumors. The most prevalent and lethal tumor of the brain is the astrocyte-derived glioblastoma multiforme [2]. This rapidly growing recurrent cancer is resistant to all conventional treatments. Moreover, malignancies of the brain are difficult to manage because most agents cannot cross the blood-brain barrier and the healthy part of the brain is very susceptible to damage as a result of therapeutic interventions [3]. In research, 3D neurosphere culture of brain cancer cells is considered a biologically relevant in vitro model of the zones of the tumor that exhibit the highest level of malignancy [5]. The neurosphere culture system was first described by Reynolds and Weiss 1992 [9] and involves the proliferative cultivation of neural stem cells and their progeny as free-floating spherical clusters, so-called neurospheres. In the neurosphere culture system using neural cancer cells, neurospheres presumably develop from individual cancer stem cells (CSC) while closely reflecting and maintaining the genetic and phenotypic traits of the primary tumor [6, 7]. Neurosphere-derived cells express stemness markers that are typical of neural stem cells (NSC), e.g. Nestin and Sox2, but usually lack the differentiation marker GFAP [1, 5].

Brain tumor CSCs, which share multiple properties and hallmarks of NSCs, are a small subset of self-renewing tumorigenic cells that divide fairly slowly [4]. Similarly to the CSC subpopulations contained in the primary tumor, they are multipotent and possess significant differentiation potential for generating cell types of the neural lineage in vitro [5-8].

Adequate culture systems that induce efficient expansion of brain tumor cells without diminishing their self-renewal capabilities or differentiation potential are an essential prerequisite for research aimed at identifying ways of eradicating brain tumor CSCs with targeted, holistically curative therapies.

Schematic Overview

![Schematic Overview](image)

**Fig. 2:** Schematic overview of the origin, evolution and fate of cancer stem cells (CSC).
CSC Neurosphere Culture Protocol

I. Materials

- Cancer Stem Cell Medium (C-28070)
- Phosphate Buffered Saline w/o Ca++/Mg++ (PBS, C-40232)
- Detach-Kit (C-41210)
- 6-well Suspension Culture Plates (e.g. Greiner Bio One, No. 657 185)
- Cultured neuronal cancer cell line (for initial neurosphere culture set-up)

II. Initiation Protocol for CSC Neurosphere Cultures

1. Harvest the adherent cells
   The neural cell culture to be used for the setup of neurospheres should be in good condition. Harvest the neural cells as a single cell suspension according to your standard procedure. Resuspend the cells in a small volume, e.g. 3 – 5 ml, of the Cancer Stem Cell Medium.

2. Count the cells
   Count the cells using your routine method and adjust the volume with Cancer Stem Cell Medium to obtain a concentration of 1 million cells/ml.

3. Set up the viable culture
   Seed the cells in appropriate suspension culture vessels at 10,000 cells/ml, e.g. 40,000 cells in 4 ml of Cancer Stem Cell Medium in each well of a 6-well suspension culture plate.

4. Allow the neurospheres to grow
   Incubate the culture for 4 – 10 days, depending on the cell type used. Add one half of the culture volume of fresh Cancer Stem Cell Medium every 3 – 4 days. Do not change the medium.

5. Passage of the neurosphere culture
   The neurospheres should be passaged (section III. below) before they start to develop a dark center. Depending on the cell type used optimal passage should occur after 4 – 10 days.

III. Serial Passage of CSC Neurosphere Cultures

1. Collect the neurospheres
   Transfer the Cancer Stem Cell Medium containing the neurospheres into 15 ml conical tubes using a serological pipet.

2. Gravity sedimentation of the neurospheres
   Allow the spheres to settle by gravity sedimentation for 10 – 12 minutes at room temperature. Aspirate the supernatant, but leave approximately 200 μl in the conical tube. Do not aspirate the neurospheres.

3. Wash the neurospheres
   Repeat the sedimentation (step 2) with an equal volume of PBS. Gently aspirate the PBS leaving approximately 200 μl in the conical tube.
4. **Enzymatic digestion of the neurospheres**

Add 1 ml of Trypsin-EDTA to the neurospheres and incubate for 2–4 minutes at room temperature. Keep the spheres resuspended in the trypsin solution by pipetting up and down once every 30 seconds. Avoid sedimentation of the spheres.

**Note:** The optimal incubation time required to achieve complete dissociation in the following step III. 5 (below) must be determined empirically by the user for each cell type. While 2–3 minutes will be optimal in most cases, neurospheres of some cell lines may need longer incubation. If a completely defined dissociation process is preferred, a recombinant trypsin solution may be used as an alternative dissociation reagent according to the supplier’s instructions.

5. **Break down remaining cell aggregates**

Pipet the spheres up and down 10–20 times using a 1000 μl pipet tip to generate a single cell suspension. Aspirate the cell suspension as normal but tilt the pipet tip slightly at the bottom of the tube when expelling the cells. The shear forces generated facilitate the break-up of any residual cell aggregates. Perform a visual check to confirm that no large cell aggregates remain. Immediately after trituration, add twice the volume of Trypsin Neutralization Solution (TNS).

**Note:** Do not over-triturate as cell viability will be compromised. If in doubt, monitor the dissociation process microscopically. Non-dissociated cell aggregates may be removed by passing the cell suspension through a 40 μm cell strainer. When using recombinant trypsin use fresh Cancer Stem Cell Medium for inactivation instead of TNS.

6. **Determine the cell number and viability**

Make up to 5 ml with fresh Cancer Stem Cell Medium and determine the cell number and viability. Centrifuge the cells for 5 minutes at 300 x g. Discard the supernatant and resuspend the cells in fresh Cancer Stem Cell Medium at 1 million cells/ml.

**Note:** Alternatively, the cells may be resuspended in buffer, e.g. PBS w/o Ca++/Mg++ plus 0.5% albumin plus 2 mM EDTA, and used for further experiments and/or analytical procedures.

7. **Plate the cells**

Reseed the cells at 10,000 cells/ml in new suspension culture vessels. Typically, 6-well plates with 40,000 cells in 4 ml of medium per well are used.
Supplementary Data

Fig. 3: U-87 MG cells cultured as 3D neurospheres (passage 1 and 10). In passage 1, the competing medium (upper right) exhibits neurosphere formation comparable to that of the Cancer Stem Cell Medium (upper left). However, the culture in Cancer Stem Cell Medium results in significantly more cell mass already in this early passage (upper left). In serial passage 10, the competing medium produces slowly proliferating shapeless cell clusters (lower right), while the Cancer Stem Cell Medium consistently results in uniform, quickly growing neurospheres (all images are at 100x magnification).
Products

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Related Products

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<td>DetachKit</td>
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References

[1] Lee, J., et al., Tumor stem cells derived from glioblastomas cultured in bFGF and EGF more closely mirror the phenotype and genotype of primary tumors than do serum-cultured cell lines. *Cancer Cell*, 2006. 9(9): 391-403.


