Effectiveness of gelatin solidification for unfrozen preservation and transportation of astrocytes

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ABSTRACT

We herein studied a non-cryogenic cell preservation and transportation method. We found that astrocytes can be preserved in an unfrozen state at 4°C (without medium exchange), while maintaining good condition in comparison with cryopreservation for the periods of 5 - 7 days. The gelatin solidification method can also prevent decreases in survival rate by preventing damage to astrocytes during transportation. Therefore, the gelatin solidification method at 4°C may be a useful choice for short-term preservation and transportation.

Keywords: Cell Preservation; Gelatin; Astrocytes; Unfrozen Preservation; Transportation

1. INTRODUCTION

Cryopreservation at or near the temperature of liquid nitrogen, is commonly used in biological research. However, freezing induces cell death due to intracellular ice formation [1-5] or chemoosmotic stress [3,6-10], which results in plasma membrane disruption and subsequent necrosis. To prevent ice formation, numerous investigators have cryopreserved cellular systems using cryoprotective agents (i.e., dimethyl sulfoxide, glycerol or sugar) in an extracellular-like carrier solution, such as standard cell culture medium, but this approach is associated with harmful effects due to the cryoprotective reagents themselves [11,12]. Therefore, it is desirable that cells be preserved in an unfrozen state.

When cells are transported in culture medium in an unfrozen state, they are subjected to physical stress and damage. Therefore, cell solidification is important for transportation in such a state. Agar is widely used for cultivation and is considered to be harmless to cells; however, the melting temperature of agar is above 60°C, and cells can be damaged at this temperature. Thus, we attempted to use gelatin, which solidifies at about 15°C - 20°C. Gelatin is known to be safe for cells, as gelatin-coated dishes are routinely used for cell culture, and gelatin was used as a solid medium before agar [13,14].

For this experiment, we selected astrocytes, which are a sub-type of the central nervous glial cells, and are the most abundant cell in the human brain. They perform numerous functions, including biochemical support of endothelial cells that form the blood-brain barrier, provision of nutrients to the nervous tissue, maintenance of extracellular ion balance, and acting in the repair and scarring process of the brain and spinal cord following traumatic injuries [15,16].

We studied the survival rate of these cells in an unfrozen state for up to one week, which would be necessary for domestic transport in Japan. We first compared the survival rate and metabolic activity of astrocytes between unfrozen preservation and cryopreservation. We then determined the optimal conditions (gelatin concentration, dissolution temperature, solidification temperature, etc.) for the gelatin solution, and investigated the influence of gelatin solidification on astrocyte survival. Finally, from the standpoint of transportation, the influence of physical stress on survival rates of astrocytes preserved under two different conditions (gelatin solidification and in solution) was investigated. Based on these results, we discuss the effects of gelatin solidification on survivability and protection from damage during transportation of astrocytes in an unfrozen state.

2. MATERIAL AND METHODS

2.1. Materials

Astrocytes were obtained from a neonatal rat brain-derived primary culture (CLEA Japan, Inc.). Trypsin and calf serum were obtained from Gibco BRL. Dulbecco’s modified eagle’s medium (DMEM) and phosphate buffered saline (PBS) were purchased from Invitrogen. Gelatin from porcine skin type A and trypan blue stain were obtained from Sigma-Aldrich and Gibco, respectively. The CellTiter
96® AQueousOne Solution Cell Proliferation (MTS) assay kit and sodium dihydrogen phosphate dehydrate were purchased from Promega KK and Wako Pure Chemical Industries Ltd., respectively.

2.2. Sample Preparation

Astrocytes in DMEM containing 10% FBS (fetal calf serum; GIBCO BRL) were incubated in a CO₂ incubator for 20 - 30 days at 37°C under 5% CO₂/95% air. Viable astrocytes adhering to the surface of the culture flask were detached with 0.05% trypsin (GIBCO BRL). Detached astrocytes were centrifuged at 800 rpm for 7 min, and were suspended in DMEM. Suspended astrocytes were subjected to filtration (100-μm mesh), and the number of astrocytes was counted under a microscope. The number of astrocytes used in this experiment was controlled to 1 × 10^6 in 300 μl DMEM. This 300-μl cell suspension was then placed into a 1.5-ml microcentrifuge tube and preserved for 4 days at 4°C.

10% gelatin stock solution was prepared by dissolving gelatin in PBS, and it was autoclaved 20 min at 120°C. Gelatin solutions of various concentrations were prepared by dilution in DMEM at 35°C. In this experiment, “in solution” refers to the experiment in which DMEM medium contained no gelatin. All of the added gelatin was dissolved in DMEM medium; thus, 1.5% gelatin refers to DMEM medium containing 1.5% gelatin.

2.3. Astrocyte Preservation

Astrocytes in solidified gelatin and in solution were preserved in a static state and under agitation for 4 days at 4°C. To simulate transportation, cells were agitated at 20 rpm during the preservation period using a rotator (RT-30mini; TAITEC) at 4°C. As shown in previous section the number of astrocytes was controlled to 1 × 10^6 in 300 μl DMEM, and then preserved in a 1.5-ml microcentrifuge tube.

Cryopreserved astrocytes were prepared by adding 5 × 10^5 astrocytes to 1.0 mL of preservation solution (Cell Banker). The cell suspension in a cryotube was placed on ice for 30 min, followed by cooling to −80°C and preservation for 2, 3, 4, 7 or 14 days. Cryopreserved astrocytes were quickly thawed in warm water at 35°C before use.

2.4. Determination of Survival Rates and Metabolism

After preservation under each condition, the number of viable cells was determined using the trypan blue staining method, which indicates membrane integrity; when the cell membrane is damaged, the dye penetrates through the membrane and dead cells are stained blue. The metabolic activity of cells was determined by MTS assay, a colorimetric assay for measuring the activity of enzymes that reduce MTS to formazan dyes, which have a purple color. After 4 days of preservation, each sample was incubated at 35°C in a constant-temperature water bath for 15 min in order to melt the gelatin (samples with no gelatin added were also incubated for comparison under the same conditions). In viable cells, MTS, which has a maximum absorbance at 490 nm, is metabolized to formazan by succinate-tetrazolium reductase, leading to a decrease in absorbance at 490 nm. From this change in absorbance, metabolic activity was determined.

2.5. Morphological Observation

As astrocytes are adherent cells, viable astrocytes adhere to culture plates and develop an increasing number of projections. In contrast, dead astrocytes do not adhere to culture plates and are suspended (spherical form) in medium. Therefore, the life and death of astrocytes can be qualitatively determined based on their shape in culture.

3. RESULTS

3.1. Optimal Preservation Temperature for Astrocytes in Closed Vessel

The optimal temperature for astrocyte survival in a closed vessel was determined, because during transport, we are unable to use a CO₂ incubator or exchange medium. We performed experiments at three temperatures; 4°C (refrigeration temperature), 25°C (room temperature) and 37°C (incubation temperature). Figure 1 shows the survival rates and metabolic activities of astrocytes after 4 days of preservation. Significant viability was only observed at 4°C, while the metabolic activity at 4°C was similar to that before preservation. Furthermore, astrocytes preserved at 4°C for 4 days adhered to the culture plate and showed an increasing number of projections (data not shown). These results confirm that 4°C is adequate for astrocyte preservation without medium exchange for 4 days or more. All subsequent unfrozen preservation experiments were thus performed at 4°C.

3.2. Influence of Freezing on Survival Rate and Metabolic Activity of Astrocytes

Survival rates of astrocytes after cryopreservation and unfrozen preservation are shown in Figure 2.

Cryo-preserved astrocytes at −80°C showed survival rates of approximately 80% up to 14 days. On the other hand, the survival rate of astrocytes preserved at 4°C decreased gradually with increasing preservation period. Therefore, cryopreservation becomes advantageous with longer preservation periods, as judged by the trypan blue method. In contrast, the metabolic activity of cryo-preserved astrocytes, as determined by MTS assay, decreased to
about 50% at day 14, regardless of preservation period (Figure 3), while that of astrocytes preserved at 4°C was over 70% up to day 7 (Figure 3).

3.3. pH Changes of DMEM Medium after Addition of Gelatin and Influence on Survival Rate of Astrocytes

The pH of DMEM medium decreased from 8.3 to 7.0 by addition of gelatin from 0% to 5% at room temperature (gelatin is not solidified at this temperature). Therefore, we measured the pH dependence of astrocyte survival rate in DMEM medium. The pH (6.8 ~ 8.4) was adjusted with H$_2$NaPO$_4$ buffer, and the survival rate of astrocytes showed little change. Thus, we ignored the influence of pH changes on survival rate under the present experiment condition, and used gelatin-DMEM solution without adjusting pH.

3.4. Adequate Gelatin Concentration for Astrocyte Solidification

In order to solidify the cells in an adequate period at 4°C, optimal gelatin conditions were determined. First, we measured the sedimentation time of astrocytes in 1.5-ml microcentrifuge tubes at four gelatin concentrations (0%, 0.5%, 1.5%, 3.0% and 5.0%) at 25°C. The sedimentation time was 25 min at 1.5% and 3.0% gelatin concentrations at 25°C, and these gelatin solutions were solidified with sufficient hardness within 15 min at 4°C. However, the sedimentation time of astrocytes in 5.0% gelatin solution at 25°C was over 60 min, and the solidification time in 0.5% gelatin solution at 4°C was more than 300 min. Therefore, 1.5% and 3.0% gelatin solutions were used in subsequent experiments.

3.5. Influence of Gelatin Concentration on Survival Rate of Astrocytes in Standing Preservation

Survival rates of astrocytes solidified at various gelatin concentrations (0%, 1.5% and 3.0%) were determined after standing preservation for 4 days at 4°C in order to clarify the effects of gelatin solidification. For astrocytes preserved in solution and those solidified in 1.5% gelatin, the survival rate was about 65%, however, the survival rate of astrocytes in 3% gelatin was about 10% lower.
Based on the results for sedimentation time, solidification time and influence on survival rate, we studied the effects of physical impact during transportation on survival rates of astrocytes in solution (DMEM medium) and solidified in 1.5% gelatin-DMEM medium.

3.6. Influence of Physical Impact (Agitation by Rollover Stirring) on Survival Rates of Astrocytes Preserved in Solidified Gelatin and Those in Solution

The effects of physical impact (agitation by rollover stirring) on the survival rate of astrocytes preserved in solidified gelatin was compared with that of astrocytes preserved in solution at 4°C for 4 days (Figure 4).

After 4 days, the survival rate of astrocytes (5%) subjected to physical impact was about 50% lower when compared to those preserved in standing solution (54%). On the other hand, in astrocytes preserved in solidified with gelatin, the survival rate was unchanged. In solution, physical impact dispersed and disrupted astrocytes in the preservation medium, and morphology was affected in about 75% of cells. However, astrocytes solidified in gelatin were fixed at the bottom of vessel and were not injured at all.

3.7. Influence of Physical Impact on Metabolic and Growth Activity of Astrocytes

The absorbance (MTS assay) under two conditions (solidified in 1.5% gelatin and dispersed in solution) after 4 days of standing preservation at 4°C was consistent. This indicates that there are no differences in the metabolic activity of viable astrocytes under these preservation conditions. Growth activity also showed similar values, irrespective of preservation method (data not shown). Therefore, the metabolic and growth activity for viable astrocyte after preservation by various methods showed no significant differences.

4. DISCUSSION

The survival rate of astrocytes in a standing state without medium exchange at 4°C was clearly higher than at 25°C and 37°C. The survival rate of cryopreserved astrocytes was higher than after unfrozen preservation at 4°C for preservation periods of up to one week. Clear differences were observed after 5 or 6 days of preservation. However, the metabolic activity of cryopreserved astrocytes was lower than that of astrocytes preserved at 4°C. This indicates that some of the cryopreserved astrocytes judged as viable cells had lost metabolic activity, or that the metabolic activity of all of viable astrocytes was decreased by freezing. Therefore, we compared the “metabolic activity ratio”, which is defined as the survival rate multiplied by the metabolic activity. Here, metabolic activity (normalized against the same living cell number) is calculated by dividing the absorbance at 490 nm observed on MTS assay before preservation by that after preservation. Figure 5 shows the metabolic activity ratio of astrocytes determined by above method after preservation at –80°C and at 4°C, and it was revealed that astrocytes preserved at 4°C maintained good condition (high metabolic activity ratio) when compared with cryopreserved cells over periods of 4 or 5 days.

Figure 4. Survival rate of astrocytes after 4 days of preservation at 4°C in solution and solidified in gelatin. A: Preserved under static conditions; B: Preserved with agitation by rollover stirring.

Figure 5. Preservation period dependence of metabolic activity rates of refrigerated (4°C, ○) and cryopreserved (–80°C, □) astrocytes. Here, metabolic activity is normalized against survival rate, which was determined by absorbance at 490 nm on MTS assay. Mean ± SEM (n = 3).
On the other hand, when dispersed in solution, cells are disrupted due to movement during transport. Thus, we solidified the astrocytes in gelatin. Both the melting temperature and solidification time of 1.5% gelatin were adequate for astrocyte preservation, because 1.5% gelatin solution solidified in approximately 15 min at 4°C and melted in approximately 5 min at about 37°C. When astrocytes were preserved for 4 days in solidified gelatin, the survival rate after physical impact was the same as those preserved in the standing state. In contrast, when astrocytes were preserved in solution, the survival rate in the standing state was 54%, and this decreased to about 5% after physical impact. Similar results were seen for metabolic activity.

Gelatin solidification has no risk of harmful effects, in contrast to cryoprotective agents, and there is no risk of injuries due to ice crystals [1-5,11,12]. In fact, metabolic activity rate was lower with cryopreservation (~80°C), which is a typical preservation method for periods of up to about five days, as compared with the gelatin solidification method at 4°C. These results confirm that astrocytes can be preserved in an unfrozen state at 4°C (without medium exchange) while maintaining good condition in comparison with cryopreservation for periods of 5 - 7 days. In addition, gelatin solidification is inexpensive, simple and effective for preventing damage to astrocytes during transportation. Therefore, gelatin solidification at 4°C is a useful option for short-term cell preservation and transportation.

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REFERENCES


