TITLE

A Drop Array Culture for Patterning Adherent Mouse Embryonic Stem Cell-Derived Neurospheres

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ABSTRACT

New therapeutic approaches for repairing an injured or degenerating nervous system have accelerated the development of methods to efficiently generate populations of neurons derived from various stem cell sources. Many of these methods require the generation of neurospheres. Here we describe a simple technique for creating an array of adherent mouse embryonic stem cell (mESC)-derived neurospheres using a conventional plastic culture dish and a patterning template. mESC-derived neurospheres are confined to circular (4 mm-diameter), gel-coated regions within an array. The adherent neurosphere arrays require three days to prepare from an mESC source; they can be maintained in 15 µL drops of medium, and exhibit extensive neurite elaboration after eight days of cultivation. Additionally, we demonstrate the potential of treating the adherent neurospheres in selected drops of an array with a variety of differentiation-inducing reagents and/ subsequently individually analyzing such neurospheres for gene expression, protein levels and morphological development.

KEYWORDS

Pattern, Array, Neurospheres, Drop-Culture, Stem Cells, Hanging Drops
SHORT COMMUNICATION

Self-renewal and tissue-specific differentiation are signature traits of stem cells that researchers have capitalized upon to devise cell-based therapies for the treatment of an assortment of neurodegenerative disorders including amyotrophic lateral sclerosis, Parkinson’s disease, and age-related macular degeneration (Steward et al., 2013). The promise of stem cells for replacing damaged or diseased cells in the nervous system has spurred an ongoing refinement of methods to efficiently generate homogeneous populations of “neurons” from embryonic, multipotent, or induced pluripotent stem cell sources (Torisawa et al., 2007, Steward et al., 2013).

Some methods for obtaining populations of embryonic stem cell (ESC)-derived neurons require the formation of embryoid bodies, or stem cell aggregates, which are grown with or without retinoic acid (RA). Since RA directs stem cells towards a neuroectoderm lineage (Ying et al., 2003), spheroids grown in a medium devoid of RA must be further propagated in a serum-free defined neuronal differentiation medium, which discourages growth of non-neural cells (Stavridis and Smith, 2003). Neurons can be derived from these primed spheroids (Torisawa et al. 2007), from dissociated spheroids, or directly from a monolayer of ESC under medium conditions conducive to neuronal differentiation. Other procedural elements, such as substrate coating, defined medium components, and longevity of culture can greatly vary (O'Shea, 2001).

Researchers have demonstrated that neurospheres can be cultured in an array format to create high-throughput assays (Shofuda et al., 2013) and even complex neural networks for implantation (Kato-Negishi et al., 2010). Traditionally, round bottom 96-well plates and hanging drops are the
standard platforms for generating neurospheres. We have used microfluidic approaches in the past to efficiently engineer uniform neurospheres (Torisawa et al., 2007). Shofuda et al. custom-designed a polymethylmethacrylate microchip array, dubbed microsphere array (MSA), to gain precise control over neurosphere homogeneity and enable bulk production. Neurosphere size can be controlled with different sized wells, and a polyethylene glycol (PEG) surface coating permits ease of neurosphere retrieval by hindering cell adhesion (Shofuda et al., 2013). Alternatively, a PEG-based hydrogel microwell array can be used to form neurospheres, with each neurosphere arising from a single founding cell (Cordey et al., 2008). Additionally, polydimethylsiloxane (PDMS) microchambers have been used to culture grid-like arrays of neurospheres that are interconnected with neuronal processes, and the flexible chambers have been used to “stamp” intact neuronal networks directly onto damaged brain regions in rats (Kato-Negishi et al., 2010).

Here we present a quick, simple, cheap, and efficient method for patterning an array of adherent neurospheroids on tissue culture dishes. The spheroids can be maintained in individual drops or in a flood of medium. Prior to creating this array, a single cell suspension of mouse embryonic stem cells (mESCs, D3 line) (Doetschman et al., 1985) are suspended in neuronal differentiation medium (50:50 mix DMEM/F12 and Neurobasal Medium, supplemented with 1% B27, 0.5% N2, 1% sodium pyruvate, and 1% antibiotic-antimycotic) for 24 h. During this time, the mESCs are primed for differentiation through removal of trace amounts of leukemia inhibitory factor (LIF) that preserved self-renewal and pluripotency, and exposure to a neuronal
differentiation promoting medium. This neuronal differentiation medium (NDM) is enriched in B27 and N2 (Lorincz, 2006), which have been demonstrated to be effective in guiding the early stages of differentiation of mESCs into neurons. The resulting mESCs heterogeneously-sized aggregates are subsequently dissociated into another single cell suspension that is used to create hanging drops for the formation of more uniformly sized spheroids.

The general method for forming arrays of adherent neurospheres involves the transfer of neurospheres from hanging drops onto adhesive regions already configured in a 4 x 4 circle-array on a 35 mm plastic Petri dish. Geltrex, an adhesive and differentiation-promoting agent, guides the placement of the spheres. Geltrex is a commercially available approximation of extracellular matrix, which contains laminin, collagen IV, entactin, and heparin sulfate proteoglycan (Loring and Peterson, 2012). Matrigel, an earlier developed version of this matrix, has been shown to promote ESC-derived neuron progenitor survival and processes elaboration (Uemura et al, 2010). A schematic of the adherent neurospheroid array creation process is shown in Figure 1A. Two variations of our method exist, and we refer to them as “transfer” and “inversion.”

In the “transfer” version of this method, the single cell suspension, obtained from dissociation of the primed free-floating mESC clusters, is used to form hanging drops (500 cells/10 µL) on the lid of Petri dishes. Sterile distilled water is added to each dish’s base to inhibit evaporation from the drops, and the spheres are allowed to develop for 24 h. Concurrently, arrays of Geltrex are created on the bases of a separate set of dishes (gel arrays incubated at 37°C for 24 h). Each drop of medium, containing a spheroid, is individually
removed from the lid and reformed on a circular gel-coated region of the prepared gel-array dish. Minute amounts of medium can be added to each drop to achieve a total drop volume of 15 µL.

An array template (Figure 1B), printed on paper, is positioned beneath a culture dish to guide the placement of gel or spheres as well as the removal or addition of solutions.

In the simplified “inversion” variation of this method, drops of a single-cell suspension are formed directly on gel-coated regions that are pre-pattered in an array on the dish base. The dish base is then inverted so that it rests over small silicone posts, positioned along the inner rim of the dish lid as illustrated in Figure 1A (Step 2), to ensure sufficient ventilation. The spheres formed in these hanging drops are suspended from, but not touching a gel pattern. After 24 hours, the dish is smoothly inverted so that all of the spheres make contact with the gel at once, and this contact is prerequisite for sphere-surface adhesion. This quick inversion process circumvents the time-consuming task of using a pipette to transfer each drop, one-by-one, to a single gel-coated region.

For both methods, we found the spheroids adhered firmly to the gel patterns after 24 h, as evidenced by the outward spread of cells (Figure 1D). At this point, designated day 0 for the differentiation of adherent spheroids, the cell masses can be maintained by either adding fresh drops of NDM to spheroids or gently flooding the entire spheroid array with fresh NDM. Spheroids were maintained on gel in either “drops” or a “pool” of medium for a period of 8 days (Figure 1A, Step 3, drop culture left, pool culture right). A portion of the medium was removed and replaced every other day. Specifically, in the “pool” culture, around half of the medium is
replaced with fresh NDM medium. In the “drop” culture, 7.5 µL of medium was carefully removed from each drop and replaced with 12.5 µL of fresh NDM medium. In some instances, medium in drops was refreshed daily. Neurite process formation could be observed as early as day 2, and extensive neurite outgrowth was visible at day 8 (Figure 1D).

We categorized the technique used to form adherent neurospheroid arrays as “transfer” or “inversion,” and the maintenance of cultures by the “drop” or “pool,” method. In the figures, “d” denotes daily medium exchange. The methods compared and evaluated were: transfer-drop, transfer-drop(d), transfer-pool, and invert-pool. No inversion-drop conditions were evaluated.

Cell density and hanging drop volume were optimized to rapidly generate uniform spheres (Figure 1C). In the “transfer” method, the average diameter of spheres obtained from hanging drops was 166 µm ± 15 µm (n = 80). In the “inversion” method, an average diameter 134 µm ± 22 µm (n = 153) was found. It is possible that the presence of the gel could affect the sphere formation process. Each user would need to carefully optimize hanging drop formation conditions for a chosen array culture technique.

Here we use a 16-drop template for a 35 mm-diameter dish, but some researchers may desire to form spheres on surfaces with larger areas. An array template could also be designed to match the well geometries and arrangements that are characteristic of common well-plate schemes that are compatible with liquid handling robots (Tung et al, 2011).
The region of elaborate neurite outgrowth is clearly outlined by labelling cells/processes for neuronal beta tubulin III, TUJ1 (Figure 2A). We used a morphometric analysis technique to characterize process formation of stained cell masses that were fixed at two-day intervals over a total of eight days, when using the “transfer-pool” method (Figure 2B) on day 8 for all types of culture methods (Figure 2C). The area of neurite outgrowth was obtained by subtracting the area of a nuclei cluster from the area of an entire cell mass (Figure 2D). FIJI (NIH), an image processing software, was used to obtain area measurements. Clear formation of intricate neurite elaboration is evident at day 6, where there is a substantial increase in neurite outgrowth area, compared to day 2 and 4 (Figure 2E). There was no significant difference found among the average values of neurite outgrowth areas among the four different culture methods evaluated (Figure 2E).

To confirm neural differentiation, we performed quantitative reverse transcription PCR (RT-qPCR) on cDNA made from RNA isolated from cells in one-day-old spheroids suspended in hanging drops at day 0’, and also those adhered to Geltrix patterns at day 4 and day 8 (Figure 2F). The “transfer-pool” group of cells was used to harvest RNA from day 4 and day 8 spheroids. We screened our neurospheroids for the temporal expression of the neuron-specific markers, beta tubulin III (Tubb3) and synaptophysin (Syp), the embryonic stem cell marker, octamer-binding transcription factor 4 (Oct-4), and the glial cell marker, glial fibrillary acidic protein (GFAP). The 8-day-old spheroids expressed significantly higher levels of Tubb3 and
Syp and lower levels of Oct-4 and GFAP than day 4 and day 0’ spheroids. These results suggest that the spheroids are directed toward at least the early stages of neuronal development.

To demonstrate the usability of this method for treating spheres in an array selectively, we treated drops of cells with green fluorescent cell tracker according to a specific configuration (Figure 2G). Ideally, a neurosphere array representing different brain regions could be constructed by including neurospheres, derived from selected brain regions, within different drops at specific locations on the array. These regions could be homogenously or selectively treated with various agents to assess therapeutic or neurotoxic effects. 1,3-Dinitrobenzene is an example of one neurotoxicant that causes brain-region specific damage (Dixon and Philbert, 2015).

The distances between the spheroids could be optimized for specific biological assays. The effects of longer-duration cultures (>1 week) and variable interspacing lengths on neurospheroid development need further experimentation.

The novelty of this system lies in its multipurpose features that include the formation of adherent neurospheroids arrays inside drops of solution, maintenance of adherent neurospheroids inside drops or a pool of medium for several days, and the use of this method to pattern the treatment of adherent neurospheroids, selectively. This simple, yet versatile, system for creating adherent spheroid arrays and drop cultures could prove useful not only for neuroscience studies, but also other research areas where prerequisite formation of uniform adherent spheres is required.
A detailed procedure is supplied as supporting information.

The authors declare that there is no conflict of interest.

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A.R.D. designed and performed experiments, analyzed the data, and wrote the manuscript. Y.R. performed experiments and wrote the manuscript. K.H. performed experiments, analyzed the data, and wrote the manuscript. K.F.B. wrote and edited the manuscript.

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FIGURES AND LEGENDS
Figure 1. In Dish Drop Array Culture of Neurospheres. A. Step 1. Geltrex drops (10μL) are arranged on the base of an untreated dish and incubated for 24 h at 37°C. Step 2. In the “transfer” method, drops from a suspension of primed mESCs are created on dish lids, and the dish (10cm-dia.) is filled with water (0.1 mL/cm²) to curb evaporation. In the “inversion” method, hanging drops are formed directly on gel regions covering the base of the dish. A well and three posts, all machined from PDMS slabs, were sealed respectively at the center and along the perimeter of the Petri dish. The dish containing drops was inverted over the well to rest on
the posts and over a well filled with water (~0.5mL). **Step 3.** After 24 h, spheres from both methods are placed in contact with the gel-coated regions pattern. For the “transfer” method, a pipette is used to transfer a sphere-containing drop to a gel coated region of a dish pre-patterned with a gel array. In the “inversion” method, wells and posts were removed from the dish lid, and the set-up was inverted, allowing spheres in the array to make contact the existing gel. Spheres in both methods were allowed to firmly attach to the gel array for 24 h., after which, the cells masses can be maintained in “drop” or “pool” culture formats. **B.** A template with a 4 x 4 array of 4mm-diameter circles was created for a 35mm-diameter Petri-dish. The template is placed beneath the dish (1), used to guide deposition of drops (2), and then removed following the completion of drop formation (3). This template also serves as a place- holder when it is realigned beneath gel drops, prior to their removal, and during subsequent formation of drops containing cells. **C.** After 24 h, fully formed spheres can be observed inside of the hanging drops that were initially composed of 500 cells in 10µL neuronal differentiation medium (5x10^4 cells/mL). **D.** The cell masses were cultured for 8 days. Day 0 was designated as the day the spheres adhered to the gel pattern. At day 8, neurite outgrowth/process formation was clearly visible.
Figure 2. Characterization and Patterned Treatment of Differentiated Neurospheres. A. Fluorescent image of a complex neuritic elaboration from an 8-day old neurosphere stained with neuron-specific beta-III (TUJ1) and 4',6-diamidino-2-phenylindole (DAPI). B. Temporal expression of TUJ1 for neurospheres cultivated with the “transfer-pool” method. C. Comparison of differentiated neurospheres stained for TUJ1 at Day 8, using the indicated culturing methods. D. Quantification for the area of the neurite outgrowth was performed by
subtracting the area of a nuclei-cluster from the area of the entire cell mass. FIJI was used to create a binary image of the cell body mass, outline the nuclei-cluster, and perform area calculations. E. Neurite outgrowth areas on day 2, 4, 6 and 8 for the “transfer-pool” technique and day 8 for the indicated types of culturing techniques. Group means were determined and the analysis done using one-way ANOVA with Tukey’s post-hoc procedures. (*p-value < 0.05; error bars show standard deviation). F. Relative gene expression, compared to mESC control, of beta 3 Class III tubulin 3 (Tubb3), synaptophysin (Syp), octamer-binding transcription factor 4 (Oct-4), and glial fibrillary acidic protein (GFAP). Data was obtained by taking triplicate measurements from 3 independent RTqPCR experiments. Group means were analysis using one-way ANOVA with Tukey’s post-hoc procedures. (*p-value < 0.05; error bars show standard deviation). G. Two of the many possible patterning schemes are shown with colored drops (1). A “transfer-drop” culture was selectively treated with cell tracker on day 8 (2a & 2b). The fluorescent staining demarcates the pattern and cell morphology (2b) of the same treated drop shown in bright field (2a). Insets show the entire neurospheroid array.