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Large-Scale Production of Mature Neurons from Human Pluripotent Stem Cells in a Three-Dimensional Suspension Culture System

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Summary

Human pluripotent stem cells (hPSCs) offer a renewable source of cells that can be expanded indefinitely and differentiated into virtually any type of cell in the human body, including neurons. This opens up unprecedented possibilities to study neuronal cell and developmental biology and cellular pathology of the nervous system, provides a platform for the screening of chemical libraries that affect these processes, and offers a potential source of transplantable cells for regenerative approaches to neurological disease. However, defining protocols that permit a large number and high yield of neurons has proved difficult. We present differentiation protocols for the generation of distinct subtypes of neurons in a highly reproducible manner, with minimal experiment-to-experiment variation. These neurons form synapses with neighboring cells, exhibit spontaneous electrical activity, and respond appropriately to depolarization. hPSC-derived neurons exhibit a high degree of maturation and survive in culture for up to 4–5 months, even without astrocyte feeder layers.

Introduction

With the seminal discovery of human pluripotent stem cells (hPSCs) (<u>Thomson et al., 1998</u>, <u>Takahashi et al.,</u> 2007), human cells that would be difficult or impossible to obtain can be produced using in vitro cell-culture techniques. This in turn has raised hopes that hPSCs can be used to study and treat different forms of disease, including neurological and neuropsychiatric disorders (<u>Dolmetsch and Geschwind, 2011</u>, <u>Fox et al., 2014</u>, <u>Han et al., 2011</u>, <u>Imaizumi and Okano, 2014</u>, <u>Kanning et al., 2010</u>, <u>Liu and Zhang, 2010</u>, <u>Mariani et al., 2015</u>). However, a key step in the utilization of hPSCs for these purposes is the ability to obtain cell types of interest. This has often proved to be challenging for several reasons including neural diversity, culture-to-culture and line-to-line variability, and limitations on large-scale cell production.

Several methods have been described to obtain neurons of specific subtypes through differentiation of hPSCs, either via formation of three-dimensional (3D) embryoid bodies (EBs) or using monolayers as starting material

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(Amoroso et al., 2013, Boissart et al., 2013, Boulting et al., 2011, Eiraku and Sasai, 2012, Eiraku et al., 2008, Espuny-Camacho et al., 2013, Hu and Zhang, 2009, Kim et al., 2014, Li et al., 2009, Qu et al., 2014, Shi et al., 2012, Zeng et al., 2010). An alternative approach is transcriptional programming, whereby the forced overexpression of a cocktail of transcription factors instructs PSCs, fibroblasts, or other cell populations to adopt a specific neuronal fate (Hester et al., 2011, Vierbuchen et al., 2010). These methods have provided important insights into human neurogenesis and the pathogenesis of neurodevelopmental disorders, but they have limitations. For instance, EB-based protocols generally have comparatively low efficiencies (10%–40%) and require a relatively long time in culture to generate functional motor neurons. In addition, the neurons generated often require cellular feeder layers to survive for longer times in culture (Hu and Zhang, 2009, Boulting et al., 2011, Amoroso et al., 2013). Moreover, EB methods typically result in the formation of spheres of cells varying in size and shape, leading to differences in the kinetics and efficiency of differentiation within individual plates and from experiment to experiment. Monolaver-based protocols for the generation of both cortical and motor neurons have also been published, with recent work describing improved efficiencies (<u>Ou et al., 2014</u>). However, a disadvantage of this adherent monolayer-based protocol is that the neurons need to be passaged, and successful long-term culture after replating has not been described. Another common theme in the field has been the problem of obtaining mature cells from hPSCs. It has been shown that maintaining differentiated cells in culture can be challenging, thereby precluding experiments studying aspects of cellular functions that take longer times to manifest (Bellin et al., 2012, Grskovic et al., 2011).

Recently, a 3D culture system that yields brain tissue from hPSCs in the form of neural organoids has been described (Bershteyn and Kriegstein, 2013, Lancaster et al., 2013, Sasai, 2013). These organoids produce neurons organized in a manner reminiscent to what is seen in distinct anatomical structures within the mammalian CNS. At least some of the neurons in the organoids are functional, and this method has thereby offered a promising approach to study neurodevelopmental mechanisms and disorders. However, at this point, formation of neural organoids is not a process that is fully controlled. Another promising recent report based on a scaffold-free plate-based 3D method used to generate spheroids showed the possibility of yielding functional neurons with properties of deep and superficial cortical neurons (Pasca et al., 2015). However, this method may be difficult to implement for large-scale production of neurons and also generates cellular structures that are large enough to be potentially subject to necrosis in the core regions of the spheroids.

Here, we describe a method for large-scale production of neurons from multiple lines of human embryonic stem cells (hESCs) and human induced PSCs. We show that our method, based on the differentiation of 3D hPSC spheres maintained in suspension in spinner flasks (hereafter referred to as spin cultures), gives a higher purity and larger absolute number of cells, and has the potential to make functional neurons that can be maintained in culture for extended periods of time. Importantly, we have applied this method to the production of both cortical neurons of multiple types and spinal cord motor neurons. For both neuronal subtypes, we were able to document not only expression of appropriate marker genes but also several characteristics of mature neurons. The obtained neurons respond robustly to depolarization, form synapses as determined by punctate staining with antibodies against established synaptic proteins, and exhibit spontaneous neural activity assessed by electrophysiology and fluctuations in intracellular Ca²⁺ levels. Neurons generated from spheres also exhibited increased survival after dissociation compared with EB-derived neurons, and could be kept in culture without glial feeder layers for up to 5 months without compromising their functional properties. Taken together, these data illustrate that this protocol robustly generates large numbers of functional neurons that can reach a high degree of maturation, and should offer an improved cellular source of human neurons for disease modeling and drug screening.

Results

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Spin-Culture hPSCs Form Spheres that Adopt a Cortical Identity after Dual SMAD Inhibition

It has previously been shown that hPSCs can be cultured in 3D spheres, consisting of hundreds of individual hPSCs, in a suspension-based system that can be readily scaled to permit the production of hundreds of millions of cells per culture (Ismadi et al., 2014, Otsuji et al., 2014, Pagliuca et al., 2014). Moreover, recent work has shown that hPSCs grown under such conditions can be differentiated into pancreatic β cells that were more mature than those produced in standard two-dimensional cultures (Pagliuca et al., 2014).

To explore the potential for generating cortical neurons using suspension-based cultures, we initially employed a differentiation protocol based on inhibition of the activin/transforming growth factor β (TGF- β) and bone morphogenetic protein (BMP) signaling pathways (Chambers et al., 2009) (illustrated schematically in Figure 1 A). Prior to differentiation, the size and general appearance of hPSC spheres was quite homogeneous, and virtually all cells expressed the pluripotency markers OCT4 and NANOG (Figures S1A and S1B). During differentiation, spheres remained homogeneous in size (Figure 1B), and qRT-PCR analysis of RNA extracted from cells collected at different time points showed a rapid loss of expression of the pluripotency genes *OCT4* and *NANOG*, and gradual expression of cortical progenitor, and then neuronal genes (Figure S2). The neural progenitor genes *OTX1* and *OTX2*, and *FOXG1*, an established marker of cortical progenitor cells, reached maximum levels of expression around day 11 of differentiation, while the temporal expression profile of genes specific for mature cortical layers seemed to resemble the sequence seen during embryonic development, with *TBR1* (a specific marker of neurons in cortical layer VI) appearing first, *CTIP2* (layer V) thereafter, and *SATB2* and *CUX1* (layers II, III, and IV) at later time points (Figure S2).



Figure 1

3D Neuronal Spheres from hPSC Lines Adopt a Predominantly Cortical Identity by Default

At day 50 of differentiation, spheres were collected and processed for immunocytochemistry using antibodies for cortical and synaptic markers. The results showed that spheres express the pan-neuronal marker MAP2, cortical neuron markers TBR1, CTIP2, and SATB2, and the synaptic marker SYNAPSIN1 (Figures 1C, S4A, and S4B). Immunocytochemistry performed on sections of spheres indicated that neurons positive for markers of the different cortical layers were not arranged randomly, but rather appeared in groups expressing markers of particular cortical layers. To further substantiate this notion, we performed 3D light-sheet fluorescence microscopy (LSFM) after clearing the spheres using the SeeDB method (Ke et al., 2013), allowing us to obtain a 3D image of stained spheres. The data confirmed that neurons expressing markers for similar cortical layer identities were organized in clusters rather than being mixed in a random, stochastic manner (Figures 1D–1F and S1C; Movies S1A–S1C). To obtain results with better cellular resolution to more readily quantify the efficiency and kinetics of neuronal differentiation, we dissociated, plated, and cultured spheres differentiated for 40 days for an additional 7 or 22 days. Immunocytochemistry revealed the presence of neurons expressing the cortical markers TBR1, CTIP2, and SATB2 (Figures 2A and 2B). Unbiased quantification revealed that cultures with up to 70% of all cells expressing one of these three markers could be obtained (Figures 2C-2E; Table S1). The remaining cells were positive for the neural progenitor markers OTX1 and OTX2 and the proliferation marker Ki67, consistent with the notion that these cells are residual cortical progenitors that have not yet differentiated to post-mitotic cortical neurons. Furthermore, we found that the number of Ki67-positive cells declined with increasing time in culture, suggesting that the progenitors would ultimately be depleted from the culture (Figure S3). We did not detect any cells stained using an antibody raised against the glial marker GFAP, arguing against the presence of any astroglia in these cultures. Importantly, the spin-culture protocol facilitates the generation of large numbers of neurons. From one 300-ml spinner flask we routinely obtained 3×10^9 neurons.

Importantly, presumably because the continuous agitation provided optimal oxygenation and nutrient penetration, this spin-based method allows cells in spheres to thrive for months without any signs of having necrotic cores.



Figure 2

Neurons from Dissociated Cortical Spheres Maintain the Differentiated Phenotype in Culture

These results show that hPSC spheres in spinner flasks differentiate efficiently to neuronal subtypes with neurons acquiring a predominantly cortical identity in the absence of patterning factors. Cortical cells expressing markers of different layers were readily identified and clustered together within the 3D spheres, with CTIP2-expressing neurons predominantly residing centrally, SATB2-expressing neurons on the outside of the spheres, and TBR1-positive neurons sandwiched in between the other two populations. When dissociated and cultured as a monolayer, the neurons maintained expression of cortical markers.

Dissociated Cortical Neurons Form Spontaneously Active Neuronal Networks and Exhibit Stimulation-Induced Activity

We thereafter sought to further characterize dissociated post-mitotic neurons. We analyzed these neurons for synaptic markers using antibodies specific for MAP2, the pre-synaptic marker SYNAPSIN1, and the post-synaptic marker GLUR1, specific for glutamatergic synapses. The results showed the presence of distinct SYNAPSIN1- and GLUR1-immunoreactive puncta distributed along the MAP2-positive dendrites. As we and others have observed in monolayer cultures of hPSC-derived neurons, SYNAPSIN1 and GLUR1 immunoreactivity were not always completely overlapping, but immunoreactive puncta were frequently found in close apposition, suggesting the formation of synaptic connections (Figure 3A).



Figure 3

Dissociated Cortical Neurons Are Spontaneously Active

To seek evidence for functional synapses more directly, we asked whether hPSC-derived cortical neurons exhibit spontaneous electrical activity. We first monitored intracellular Ca^{2+} dynamics as a surrogate for electrical activity by performing time-lapse microscopy on cultures of dissociated neurons transduced with an adenoassociated virus expressing the genetically encoded Ca^{2+} sensor GCaMP6s under the control of the human synapsin promoter (AAV Syn:GCaMP6s) (<u>Chen et al., 2013</u>). Cells were transduced 3 days after dissociation and imaged every 10 days. Early after dissociation, neurons exhibited low levels of activity which increased over time, and all cultures of cortical neurons exhibited synchronized Ca^{2+} waves 60 days after dissociation (<u>Figure 3</u> B; <u>Movies S2</u> and <u>S3</u>). To further examine the electrophysiological characteristics of neurons from dissociated cortical spheres, we conducted patch-clamp recordings. After 6 weeks in culture, cortical neurons showed typical voltage-gated Na⁺ and K⁺ currents (<u>Figure 3C</u>) and were able to fire action potentials upon current injection (<u>Figure 3</u>D). Spontaneous firing of action potentials and post-synaptic currents (<u>Figures 3</u>E and 3F) was also observed in most cells recorded (4 of 5), showing that cortical neurons obtained from dissociated spheres are able to mature into functional, electrophysiologically active cells.

It is well known that synaptic activation through depolarization induces a transcriptional response in neurons (Ebert and Greenberg, 2013, Flavell et al., 2008, Hong et al., 2008, Lin et al., 2008). A key event in many types of neurons, including cortical neurons, is the phosphorylation of the transcription factor CREB. To examine

whether neurons from dissociated cortical spheres respond to depolarization in this way, we treated them with high levels of extracellular KCl. Immunostaining with a phospho-CREB-specific antibody revealed robust phosphorylation of CREB in these cultures (Figures 4A and 4B; Table S2). We then performed transcriptional analyses of KCl-stimulated cortical neurons and found substantially increased transcription of known activity-regulated genes, including the early-induced genes *NPAS4*, *cFOS*, and *ARC* by 1 hr after stimulation and the late-induced gene *BDNF* after 6 hr (Figure 4C). These results are similar to those observed using freshly isolated rodent cortical neurons (Flavell et al., 2008, Hong et al., 2008, Lin et al., 2008). Similar results were also obtained from KCl stimulation of whole intact spheres after 20 days of differentiation (Figure S4).



Figure 4

Dissociated Cortical Neurons Respond to Depolarization

Taken together, these data demonstrate that cortical neurons obtained from hPSCs cultured and differentiated as spheres in spinner flasks are electrophysiologically active and form neural networks in culture. Moreover, the neurons display voltage-gated Na⁺ and K⁺ currents, intracellular Ca²⁺ dynamics, and activity-induced gene expression characteristic of cortical neurons, further underlining their maturation into functional cortical neurons.

Motor Neuron Production from Spin-Culture hPSCs

In the next set of experiments, we explored whether the spheres would respond to exogenously supplied factors by differentiating to other neuronal subpopulations. To accomplish this, we set out to modify standard motor neuron differentiation protocols (schematic in Figure 5A). By supplying retinoic acid and sonic hedgehog (Wichterle et al., 2002), spheres rapidly differentiating into motor neurons were obtained. qRT-PCR analyses revealed a transient peak in expression of the neuronal progenitor gene *SOX1*, followed by expression of the motor neuron-specific genes *ISL1* and *HB9* (Figure S5). At later time points, increasing expression of the biosynthetic enzyme choline acetyltransferase (*ChAT*), necessary for the synthesis of acetylcholine, and the mature pan-neuronal marker *MAP2* was detected (Figure S5). When analyzed by immunocytochemistry at day 20 of differentiation, cells in spheres expressed ISL1, MAP2, ChAT, and SYNAPSIN1 (Figures 5B, S6A, and S6B).



Figure 5

3D Neuronal Spheres from Human PSC Lines Differentiate into Motor Neurons in the Presence of Patterning Factors

After 15 days of differentiation, motor neuron spheres were dissociated, plated, and cultured under feeder-free conditions for an additional week. Immunocytochemistry revealed that up to 85% of the dissociated cells expressed ISL1, with consistent results obtained in different cell lines and low experiment-to-experiment variability (Figures 5C and 5D; Table S3). We also found that the vast majority of cells in the cultures were immunopositive for ChAT (Figures 5E and 5F; Table S3). Cells from dissociated motor neuron spheres also exhibited punctate SYNAPSIN1 immunoreactivity along MAP2-positive neurites (Figure 6A). We also performed experiments in which spin-culture-derived motor neurons were cultured with human myotubes. Staining with fluorescently labeled α -bungarotoxin (BTX), a toxin that specifically binds to post-synaptic acetylcholine receptors, labeled regions of contact between motor neurons and muscle fibers (Figure 6B), indicating formation of neuromuscular junctions.



Figure 6 Dissociated Motor Neurons Are Spontaneously Active

Dissociated Motor Neurons Exhibit Stimulation-Induced Activity and Form Spontaneously Active Neural Networks

To further address the functionality of the motor neurons that we produced, we characterized their physiological properties. First, we confirmed that the cells were capable of responding to high KCl depolarization by phosphorylating CREB (Figures S6C and S6D). Next, we asked whether these motor neurons exhibited spontaneous electrical activity in culture. As before, we monitored intracellular Ca²⁺ dynamics in cultures of dissociated motor neurons transduced with AAVSyn:GCaMP6s. Cells were transduced 3 days after dissociation and imaged every week. As early as 1 week after dissociation, motor neurons already exhibited significant levels of activity, which increased over time. Neurons again displayed more synchronicity in their firing patterns at later times, suggesting the formation of neural networks, and with groups of motor neurons exhibiting synchronized Ca²⁺ dynamics 3 weeks after dissociation (Figure 6C; Movie S4).

Patch-clamp recordings were performed to investigate the electrophysiological characteristics of motor neurons from dissociated spheres. Motor neurons were found to fire action potentials spontaneously and upon current injection even after up to 3 months in culture (Figure 6D, see also Figure S7). Furthermore, GCaMP6s-mediated monitoring of intracellular Ca²⁺ dynamics showed that motor neurons responded to both γ -aminobutyric acid (GABA) and α -amino-3-hydroxy-5-methyl-4-isoxazole propionate (AMPA), as might have been expected (Figure 6E).

Thus, the sphere differentiation protocol was successfully modified to allow the generation of billions of motor neurons from one single differentiation experiment. The yield of motor neurons was very efficient, up to 85%, with little variation between individual experiments, and sphere-derived motor neurons could be cultured without glial feeder layers. Moreover, the motor neurons expressed ChAT and were physiologically active.

Increased Survival and Function of Motor Neurons Obtained through the Spin-Culture Protocol Compared with EB-Derived Motor Neurons

Throughout the course of the experiments presented, we noted a marked increase in survival of spin-culturederived neurons compared with EB-derived neurons in all assays requiring manipulations such as dissociation, fluorescence-activated cell sorting, and viral transduction. In an attempt to address this in a quantifiable manner, we generated motor neurons in parallel using two methods (but with exactly the same medium composition), namely spin-culture differentiation and EB formation in low-attachment tissue-culture plates using protocols standard in our laboratory (Ng et al., 2015, Yang et al., 2013). Motor neuron spheres in spin culture were uniform and homogeneous, whereas motor neuron differentiation using EBs led to formation of cell clumps exhibiting marked heterogeneity with respect to size, shape, and general appearance (Figures 7A, S7I, S7J, and S7K). Expression analyses by qRT-PCR of cells undergoing differentiation at different time points revealed that motor neuron spheres exhibited higher levels of expression of *ISL1*, *HB9*, and *ChAT* compared with conventional EB-derived motor neurons (Figures 7C–7E). When spheres and EBs were dissociated and neurons plated, immunocytochemistry showed that spin-culture-derived motor neurons co-expressed ISL1, MAP2, and ChAT to a higher degree than EB-derived motor neurons (Figure 7B).

Figure 7



Sphere-Derived Human Motor Neurons Survive Longer in Culture and Mai Functionality throughout Extended Time in Culture

Next, we performed a set of experiments in which we monitored the appearance and functionality of cultures of motor neurons derived from spin-culture spheres or using EB methodology. Irrespective of plating conditions, motor neurons derived from spin-culture spheres survived for a longer period of time in culture. The difference was particularly striking on coated tissue-culture plates (Figure 7F), where motor neurons from dissociated spin-culture spheres could be maintained in culture, without glial feeder layers, for several months, which in our hands proved to be impossible using motor neurons obtained through EB formation. In experiments where motor neurons were plated and transduced with genetically encoded Ca^{2+} sensors, motor neurons derived from spin-culture spheres exhibited higher amplitude and frequency in levels of fluorescence compared with EB-derived motor neurons (Figure 7G). With extended time in culture, the amplitude and frequency of fluctuations in Ca^{2+} increased in the spin-culture-derived motor neurons, whereas EB-derived motor neurons exhibited a loss in Ca^{2+} fluctuations (Figure 7G). These results are consistent with the notion that motor neurons derived from spin-culture spheres survive better and maintain their functionality during extended periods of culture.

To formally address the maturation of the different motor neuron populations, we performed comparisons using whole-cell patch recordings (Figures S7A–S7F). Our recordings of action potentials and voltage-gated Na/K currents showed that spin-culture-derived motor neurons were able to fire faster action potentials and larger Na/K currents than EB-derived motor neurons (Figures S7A–S7F). In addition, calcium imaging experiments suggest that spin-culture-derived motor neurons express functional GABA receptors and begin to undergo the transition from GABA excitation to inhibition earlier than EB-derived motor neurons (Figures S7G–S7H), which indicate a faster maturation time course (Ben-Ari, 2002). These data suggest that spin-culture-derived motor neurons may mature faster than EB-derived motor neurons with regard to their electrophysiological characteristics, although additional experiments in more hPSC lines are required to address whether this is a general phenomenon.

Discussion

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In the last 3 years, several groups have shown that neural organoids with features of the mammalian CNS can be generated (Bershteyn and Kriegstein, 2013, Lancaster et al., 2013, Sasai, 2013). This innovative approach has opened up the possibility of studying basic mechanisms of the developmental biology of the human nervous system (Camp et al., 2015, Otani et al., 2016) and neurodevelopmental disorders (Mariani et al., 2015). However, it is important to note that there are limitations to this model system. Neural organoids are produced from hPSCs through formation of neural progenitors that are aggregated individually in Matrigel and then transferred to spinner flasks for further development. The resulting organoids are highly complex structures containing many different types of neurons organized into areas exhibiting features of distinct regions of the mammalian CNS, including the cerebral cortex. However, neural differentiation in organoids is quite variable with respect to both the exact type of neuronal subtype and the structures produced, as well as the efficiency of the differentiation process.

In this paper, we present a method based on 3D growth of cells from hPSC to differentiated neurons. Importantly, this protocol does not rely on isolation and manipulation of an intermediate cell population in Matrigel. In contrast, we culture cells as spheres starting from the pluripotent state and only subject the spheres to differentiation cues by changing the composition of the culture medium. For both cortical neurons and motor

neurons produced with this method, we were able to document not only expression of appropriate marker genes but also several hallmarks of mature, functional neurons, including synapse formation, expression of functional receptors for neurotransmitters, spontaneous electrical activity, and appropriate expression of activity-induced genes.

This system has several advantages. First, it is simple to obtain a high absolute number of cells—in the range of billions even in a single flask. This is of importance for downstream applications such as high-throughput screening, where cell numbers are a limiting factor. For example, our laboratory routinely performs chemical screens on hPSC-derived neurons in 384-well plates, where 15,000 neurons are plated in each well (Yang et al., 2013). One single differentiation of hPSC-derived spheres in spinner flasks as described herein would be sufficient for approximately 350×384 -well plates, thereby greatly reducing the time and effort required to produce the cells necessary to conduct such experiments. Second, spheres cultured in the spinner flasks appear uniform in size and shape during the differentiation, reducing experiment-to-experiment variation. Third, spin-culture differentiated spheres, we were able to culture both types of neurons without the use of feeder layers for extended periods of time, as long as 4 months or more. This is in stark contrast to neurons obtained from EBs, where cells start to detach from the plate at relatively early times.

In this report we have focused on generating cortical and motor neurons, but we believe it is likely that future adaptations of the sphere protocol to generate other distinct neuronal subtypes will prove to be advantageous compared with other methods. In addition, establishing imaging and physiological assays to cells maintained in three dimensions should be possible in the near future so that functional connections among populations of neurons can be maintained and analyzed (rather than studied after dissociation). Future work will also determine whether neurons maintained in suspension for extended periods can achieve a greater degree of maturation, of major importance in the study of late-onset neurodegenerative disease. Finally, it will be important to understand whether suspension culture conditions can be adapted to allow for faster generation of neurons and glial cells. If this is the case, one of the main problems in using hPSC-derived neurons, that of prolonged times required for cell production, can be circumvented.

Experimental Procedures

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Human Pluripotent Stem Cell Lines

All experiments with the human ESC lines were reviewed and approved by the Harvard Embryonic Stem Cell Oversight Committee. The use of the wild-type iPS cell lines by the Rubin lab was reviewed by the Harvard Committee on the Use of Human Subjects (the Harvard IRB) and determined to not constitute human subjects research.

Media

mTeSR (STEMCELL Technologies); KSR medium: 15% KSR (Invitrogen), KO DMEM (Invitrogen), L-glutamine (Gibco), 100× non-essential amino acids (NEAA) (Gibco), 100× penicillin-streptomycin (Gibco), and 1000× β -mercaptoethanol (Gibco); neural induction medium (NIM): DMEM/F12 (Invitrogen), 100× N2 supplement (Gibco), 50× B27 supplement (Life Technologies) (for cortical differentiation we substituted regular B27 for B27 supplement without vitamin A [Life Technologies]), 100× Glutamax (Gibco), 100× NEAA (Gibco), 100× penicillin-streptomycin (Gibco), 125× glucose (J.T. Baker), and (only for motor neurons) 0.2 mM ascorbic acid (Sigma); neurobasal (NB) medium: Neurobasal (Invitrogen), 100× N2 supplement (Gibco), 50× B27 supplement (Life Technologies) (for cortical differentiation we substituted regular B27 for B27 supplement without vitamin A [Life Technologies]), 100× Glutamax (Gibco), 100× penicillinstreptomycin (Gibco) (for motor neurons, the medium was supplemented with 125× glucose [J.T. Baker] and

hPSC Adaption and Maintenance in Spinner Flasks

hPSCs were cultured in 500-ml disposable spinner flasks (Corning, VWR) on a nine-position stir plate (Dura-Mag) at a speed of 70 rpm, in a 37°C incubator with 5% CO₂. Prior to adaption to spinner-flask culture, hPSCs were expanded in 10-cm dishes (Corning) and dissociated to single cells with Accutase (STEMCELL Technologies) for 5 min incubation at 37°C, when they were approaching confluency. 180 million individual pluripotent stem cells were seeded into a spinner flask in 180–200 ml of mTeSR medium (STEMCELL) supplemented with 5 μM ROCK inhibitor Y-27632 (STEMCELL). Spheres formed spontaneously, and after 2 days the culture medium was changed. The cells were maintained as undifferentiated pluripotent spheres in spin culture, with medium changes every day. Every third day, when the spheres were approximately 50–100 μm in diameter, cells were passaged by incubating at 37°C for 6 min in a 1:1 mixture of Accutase and PBS, and by mechanically dissociating the cells. Cells were filtered and reseeded into the flask at the same initial density as described above. Medium was changed by taking the flasks off the stir plate, allowing the cells to settle to the bottom of the flask and the medium to be changed. At day 1 of differentiation, medium was changed to mTeSR with the activin/TGF- β inhibitor SB431542 (R&D Systems) (10 μM) and the BMP inhibitor LDN193189 (Stemgent) (1 μM) (hereafter referred to as "dual SMAD inhibition"). Subsequent steps of the differentiation protocol (as described below) use the same method for changing media.

Differentiation Conditions

Cortical Differentiation For cortical differentiation from days 2 to 10, spheres were gradually adapted to NIM through a dilution series of KSR and NIM, with dual SMAD inhibition maintained until day 7. Medium was changed as follows: days 2–4: 100% KSR medium with the Wnt signaling inhibitor XAV939 (2 μM) (Stemgent); day 5: 75% KSR medium, 25% NIM; day 7: 50% KSR medium, 50% NIM; day 9: 25% KSR medium, 75% NIM. From days 11 to 21, cultures were maintained in 100% NIM (Maroof et al., 2013). From day 21 onward, cultures were maintained in NB medium with 10 ng/ml brain-derived neurotrophic factor (BDNF) and 10 ng/ml glial cell-derived neurotrophic factor (GDNF) (both from R&D).

Motor Neuron Differentiation For motor neuron differentiation from days 2 to 10, spheres were gradually adapted to NIM through a dilution series of KSR and NIM, with dual SMAD inhibition maintained until day 6. Medium was changed as follows: day 2: 100% KSR; day 3: 100% KSR + 1 μ M retinoic acid (RA) (Sigma); day 5: 75% KSR, 25% NIM + RA, and 10 ng/ml BDNF (R&D); day 6: 50% KSR, 50% NIM + RA, 1 μ M Smoothened agonist (SAG) (Curis), and BDNF; day 8: 25% KSR, 75% NIM + RA, SAG, and BDNF. From days 10 to 15, cultures were maintained in 100% NIM + RA, SAG, BDNF, and 2.5 μ M DAPT (R&D). From day 15 onward, cultures were kept in NB + BDNF, GDNF, and 10 ng/ml ciliary neurotrophic factor (R&D).

During differentiation, spheres were collected at different time points and RNA was extracted for qRT-PCR analysis. For examination of cortical differentiation, spheres were dissociated with Accutase and plated onto Matrigel-coated plates at day 13 and analyzed the following day for expression of cortical progenitor markers by immunofluorescence. Spheres were also dissociated at day 40 with 0.05% trypsin (Gibco) and plated onto poly-D-lysine/laminin-coated (EMD Millipore/Life Technologies) plates. After 28 and 43 days of additional culture, cells were analyzed as described below. For motor neuron differentiation, spheres were dissociated with 20 units of papain per milliliter in Earle's balanced salt solution supplemented with 1 mM L-cysteine, 0.5 mM EDTA (Worthington), and DNAse. 60,000 cells/well were plated onto 96-well poly-D-lysine/laminin-coated plates at day 15 (see <u>Supplemental Experimental Procedures</u> for a detailed protocol for dissociation). Before dissociation, both cortical and motor neuron spheres were approximately 200 µm in diameter. Neurons were analyzed after one additional week in culture after dissociation.

Statistics

All experiments were performed in three independent biological replicates unless stated otherwise. A two-tailed Student's t test was used to calculate statistical significance.

Author Contributions

A.R. conceived and designed the study, carried out experiments, collected and analyzed data, and wrote the manuscript; G.R. and D.C.R. carried out experiments, and collected and analyzed data; C.S. carried out experiments, collected and analyzed data, and wrote the manuscript; F.P. contributed reagents, carried out experiments, and analyzed data; F.R., K.W., and C.B. carried out experiments; L.D. analyzed the data, Q.P.P. contributed reagents and helped with establishing spinner cultures; E.M.H. analyzed data and wrote the manuscript; L.L.R. conceived and designed the study and wrote the manuscript.

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Notes

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Footnotes

Supplemental Information includes Supplemental Experimental Procedures, seven figures, three tables, and four movies and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2016.05.010.

Supplemental Information

Document S1. Supplemental Experimental Procedures, Figures S1–S7, and Tables S1–S3:

Click here to view. (6.0M, pdf)

Movies S1A–S1C. Cortical Spin-Culture Spheres Exhibit Organization into Layers, Related to Figure 1:

Click here to view.^(284K, jpg)

Movie S2. Dissociated Cortical Neurons Exhibit Spontaneous Activity, Related to Figure 3:



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Movie S3. Dissociated Cortical Neurons Exhibit Spontaneous Activity, Related to Figure 3:

Click here to view. (390K, jpg)

Movie S4. Dissociated Motor Neurons Exhibit Spontaneous Activity, Related to Figure 5:

Click here to view. (469K, jpg)

Document S2. Article plus Supplemental Information:

Click here to view.(12M, pdf)

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