A Simple and Scalable Process for the Differentiation of Retinal Pigment Epithelium From Human Pluripotent Stem Cells

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**ABSTRACT**

Age-related macular degeneration (AMD), the leading cause of irreversible vision loss and blindness among the elderly in industrialized countries, is associated with the dysfunction and death of the retinal pigment epithelial (RPE) cells. As a result, there has been significant interest in developing RPE culture systems both to study AMD disease mechanisms and to provide substrate for possible cell-based therapies. Because of their indefinite self-renewal, human pluripotent stem cells (hPSCs) have the potential to provide an unlimited supply of RPE-like cells. However, most protocols developed to date for deriving RPE cells from hPSCs involve time- and labor-consuming manual steps, which hinder their use in biomedical applications requiring large amounts of differentiated cells. Here, we describe a simple and scalable protocol for the generation of RPE cells from hPSCs that is less labor-intensive. After amplification by clonal propagation using a myosin inhibitor, differentiation was induced in monolayers of hPSCs, and the resulting RPE cells were purified by two rounds of whole-dish single-cell passage. This approach yields highly pure populations of functional hPSC-derived RPE cells that display many characteristics of native RPE cells, including proper pigmentation and morphology, cell type-specific marker expression, polarized membrane and vascular endothelial growth factor secretion, and phagocytic activity. This work represents a step toward mass production of RPE cells from hPSCs.

**INTRODUCTION**

Age-related macular degeneration (AMD), the leading cause of irreversible vision loss and blindness among the elderly in industrialized countries, affecting 30–50 million people worldwide [1]. An early event associated with AMD is damage to retinal pigment epithelium (RPE) cells [2]. The RPE abuts the photoreceptor cell layer and has many roles in visual function, including absorption of stray light, formation of the blood-retina barrier, transport of nutrients, secretion of growth factors, isomerization of retinol, and daily clearance of shed photoreceptor outer segments [3]. RPE dysfunction and cell death is associated with both the neovascular (“wet”) and atrophic (“dry”) forms of AMD [4]. Although effective anti-vascular endothelial growth factor (VEGF)-based treatments have been developed for the neovascular form of AMD, effective treatment options for the more common atrophic form of the disease are lacking [5, 6]. One approach being explored for the treatment of atrophic AMD is the transplantation of autologous RPE [7]. However, although a potentially promising approach, harvesting autologous RPE involves complex surgery with possible sight-threatening complications [8, 9]. In addition, RPE harvested from an AMD patient, even if peripheral RPE is used as the source, runs the risk of being already impaired, because of the environmental and/or genetic factors that initially predisposed the patient to AMD.

An alternative approach to obtain human RPE cells is to generate them from human pluripotent stem cells (hPSCs), either from embryonic stem cells (hESCs) [10] or from induced pluripotent stem cells (hiPSCs) [11, 12]. Although no consensus has yet emerged regarding the equivalence of hiPSCs to hESCs [13–15], a striking feature of both cell types is their capacity for indefinite self-renewal while maintaining their potential to differentiate into all cell types of the human body. A number of investigators have demonstrated that cultured hiPSCs to hESCs can be induced to differentiate into cells that are phenotypically very similar to endogenous RPE cells. More than 10 years ago, Kawasaki et al. [16] described epithelial pigmented cells arising from differentiation of monkey ESCs on the mouse stromal cell line PA6. These cells were later
shown to express RPE markers, to phagocytose latex beads in vitro and rod outer segments in vivo, and to alleviate to some extent photoreceptor degeneration following transplantation into Royal College Surgeon rats [17]. In 2004, Klimanskaya et al. [18] generated RPE cells through spontaneous differentiation of hESCs, whereas later on, RPE monolayers were successfully differentiated from hiPSCs [19–23]. hiPSCs could thus provide an unlimited supply of RPE-like cells for transplantation in vivo [24], as well as for research applications such as disease modeling and drug studies [25]. (For the sake of language simplicity, we will from this point forward refer to RPE-like cells derived from hPSCs as “hPSC-RPE cells.”) In several animal models, hPSC-RPE cells have been shown to maintain their function after subretinal transplantation and to be able to attenuate retinal degeneration with partial preservation of visual function [17, 23, 26–29]. In addition, a human clinical trial based on hESC-RPE cell transplantation is currently under way [30].

Despite the success obtained with the generation of human RPE cells from hPSCs, published protocols to date all rely on hPSC amplification by clump passage and, following differentiation, on mechanical dissection of pigmented colonies for RPE purification [31, 32]. Both procedures, although powerful, are potentially problematic since they require time- and labor-consuming manual steps, which present challenge for scale-up and thereby constitute bottlenecks for large-scale production of high-quality and consistent hPSC-RPE cells. In this study, we describe an RPE differentiation protocol that is less work-intensive than other methods described previously. In this new protocol, hPSCs are amplified by clonal propagation using the myosin inhibitor blebbistatin [33, 34]. Subsequently, pigmented colonies are generated by spontaneous monolayer differentiation, and RPE cells are then enriched by two rounds of whole-dish single-cell passage. The procedure yields a highly purified cell population that displays many of the morphological, gene expression, and functional characteristics of native RPE cells. Completely defined conditions can be achieved through this protocol by the use of vitronectin peptide-acrylate surface (VN-PAS) [35] instead of Matrigel (BD Biosciences, San Diego, CA, http://www.bdbiosciences.com).

**Materials and Methods**

**Human Pluripotent Stem Cell Culture**

Adapting a previously described method [33], the hESC line H7 (kind gift from Dr. Hai Quan Mao, Johns Hopkins University) and the hiPSC line IMR90-4 (WiCell Research Institute, Madison, WI, http://www.wicell.org) were maintained by clonal propagation either on growth factor-reduced Matrigel or on 0.25% (wt/vol) collagenase IV (Gibco, Grand Island, NY, http://www.invitrogen.com). The loosened cell monolayer was collected after 5–10 minutes of treatment with Accutase (Sigma-Aldrich), mouse anti-ZO-1 1/500 (Invitrogen), mouse anti-BEST1 1/150 (Abcam), mouse anti-RLBP1 1/100 (Abcam), or rabbit anti-OTX2 1/500 (Millipore). The cells were then incubated for 1 night at 4°C with one of the following primary antibodies: rabbit anti- OCT4 1/500 (Abcam, Cambridge, U.K., http://www.abcam.com), rabbit anti-Nanog 1/1,000 (Millipore), rabbit anti-SOX2 1/1,000 (Millipore, Billerica, MA, http://www.millipore.com), mouse anti- SSEA4 1/100 (Millipore), mouse anti-MITF (CS) 1/100 (Thermo Fisher Scientific, Rockville, MD, http://www.thermofisher.com), mouse anti-ZO-1 1/500 (Invitrogen), mouse anti-BEST1 1/150 (Novus Biologicals, Littleton, CO, www.novusbio.com), mouse anti-RPE65 1/100 (Abcam), mouse anti-RLBP1 1/100 (Abcam), or rabbit anti-OTX2 1/500 (Millipore). The cells were then incubated for 1

**Differentiation and Culture of RPE From hPSCs**

Pluripotent stem cells were plated at 20,000 cells per cm² and maintained in TeSR1. Five days after passage, the cells formed a monolayer and were transferred to a 5% CO₂/20% O₂ incubator. Three days later, the culture medium was replaced with differentiation medium (DM) consisting of Dulbecco’s Modified Eagle’s Medium: Nutrient Mixture F-12 (DMEM/F-12) (catalog no. 11330; Invitrogen, Carlsbad, CA, http://www.invitrogen.com), 15% knockout serum (Invitrogen), 2 mM glutamine (Invitrogen), 1x nonessential amino acids (Invitrogen), 0.1 mM β-mercaptoethanol (Sigma-Aldrich), and 1x antibiotic-antimycotic (Invitrogen). Approximately 25–30 days later, pigmented foci became clearly visible and were grown for an additional 25 days. At that point, the whole monolayer of differentiating cells was passaged a first time (P1) by incubation for 4 hours in DM supplemented with 0.25% (wt/vol) collagenase IV (Gibco, Grand Island, NY, http://www.invitrogen.com). The loosened cell monolayer was thereafter broken into small clumps by vigorous pipetting. These clumps were collected by centrifugation, resuspended in Accumax (Sigma-Aldrich), and incubated for 20–30 minutes at 37°C. After vigorous pipetting, most clumps were dispersed into single cells, and the solution was filtered through a 40-µm nylon mesh (BD Falcon, San Jose, CA, http://www.bdbiosciences.com). Differentiated cells were plated at 100,000 cells per cm² on Matrigel-coated or VN-PAS plates and allowed to grow for 15–20 days in RPE medium [36] consisting of 70% DMEM (Invitrogen), 30% Ham’s F-12 Nutrient Mix (catalog no. 11765; Invitrogen), 1X B27 (Invitrogen), and 1X antibiotic-antimycotic (Invitrogen). For routine passage after P1, hPSC-RPE cells were collected by using Accumax, pelleted by centrifugation, and replated at 100,000 cells per cm².

In control experiments, hPSCs were cultured in mTeSR1 according to the manufacturer’s instructions (clump-passage method). After differentiation in DM for 50 days, the pigmented foci were isolated by manual picking as previously [37], resuspended in Accumax (Sigma-Aldrich), and incubated for 20–30 minutes in a 37°C water bath. Dissociated clumps were then directly seeded onto Matrigel-coated plates and further passaged as described above. Characterization of hPSC-RPE cells isolated through serial passage or by manual picking was performed at P2, after 50 days culture in RPE medium.

**Human Fetal RPE Culture**

Fetal RPE (fRPE) cells were obtained from ScienCell, plated at 50,000 cells per cm² on Matrigel, and cultured for 2 months in Epithelial Cell Medium (ScienCell, Carlsbad, CA, www.scienceilonline.com) before RNA was harvested.

**Immunostaining**

After fixation, the cells were blocked and permeabilized for 30 minutes in 5% goat serum, 0.25% Triton X-100 in PBS and then incubated overnight at 4°C with one of the following primary antibodies: rabbit anti-OCT4 1/500 (Abcam, Cambridge, U.K., http://www.abcam.com), rabbit anti-Nanog 1/1,000 (Millipore), rabbit anti-SOX2 1/1,000 (Millipore, Billerica, MA, http://www.millipore.com), mouse anti-SSEA4 1/100 (Millipore), mouse anti-MITF (CS) 1/100 (Thermo Fisher Scientific, Rockville, MD, http://www.thermofisher.com), mouse anti-ZO-1 1/500 (Invitrogen), mouse anti-BEST1 1/150 (Novus Biologicals, Littleton, CO, www.novusbio.com), mouse anti-RPE65 1/100 (Abcam), mouse anti-RLBP1 1/100 (Abcam), or rabbit anti-OTX2 1/500 (Millipore). The cells were then incubated for 1

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hour with the corresponding secondary antibody conjugated to Alexa 488 or Alexa 647 (Invitrogen) and counterstained with Hoechst 33342 (Invitrogen).

Mouse eye cups were fixed in 2% paraformaldehyde in 0.1 M phosphate buffer with 5% sucrose. They were then cryoprotected in 20% sucrose in 0.1 M phosphate buffer and rapidly frozen in O.C.T. compound (Tissue-Tek; Sakura Finetek, Torrance, CA, http://www.sakura.com). Eight-micrometer sections were stained with 1/250 mouse anti-rhodopsin (Lab Vision, Fremont, CA, http://www.labvision.com) conjugated to Dylight 550 according to the manufacturer’s instructions (Dylight 550 microscale antibody labeling kit; Pierce, Rockford, IL, http://www.piercenet.com) in 2% goat serum, 0.1% Triton X-100 in PBS. This was followed by incubation with Alexa Fluor 647 phallolidin (Invitrogen) and counterstaining with 4',6-diamidino-2-phenylindole.

**Western Blot Analysis**

hPSC-RPE proteins were extracted from cell lysates stored in RLT buffer (Qiagen, Hilden, Germany, http://www.qiagen.com) by acetone precipitation as previously [38] and resuspended in 1:1 vol/vol radioimmunoprecipitation (Sigma-Aldrich)-Laemmlli buffer (Bio-Rad, Hercules, CA, http://www.bio-rad.com) with EDTA-free protease inhibitor cocktail (Complete mini; Roche Applied Science, Indianapolis, IN, https://www.roche-applied-science.com). The protein concentrations were quantified using the EZQ protein quantification kit (Invitrogen). Approximately 10 μg per lane was resolved on a NuPAGE bis-tris gels (Invitrogen), and proteins were transferred to polyvinylidene difluoride membranes using the iBlot gel transfer system (Invitrogen). After a brief blocking step in SuperBlock Blocking Buffer (Thermo Scientific, Rockford, IL, http://www.thermoscientific.com) or 5% nonfat milk in Tris-buffered saline containing 0.1% Tween 20 (TBST), the membranes were incubated overnight with the following primary antibodies: rabbit anti-GAPDH 1/1,000 (Sigma-Aldrich), mouse anti-RLBP1 1/1,000 (Abcam), or mouse anti-BEST1 1/1,000 (Novus Biologicals). After several washes in TBST, the membranes were incubated in a 1/10,000 dilution of horseradish peroxidase (HRP)-conjugated anti-rabbit antibody (Santa Cruz Biotechnology Inc., Santa Cruz, CA, http://www.scbt.com) or 1:2,000 HRP-conjugated anti-mouse secondary antibodies (Cell Signaling Technology, Beverly, MA, http://www.cellsignal.com). After five washes in TBST, the membranes were incubated for 5 minutes in the SuperSignal West Femtmo Chemiluminescence Substrate (Pierce) and exposed to x-ray film (Kodak, Rochester, NY, http://www.kodak.com).

**Flow Cytometry**

For TRA-1-60, a cell surface marker, immunostaining was performed as previously [39], with a primary antibody concentration of 1 μg per 1 million cells. For intracellular and nuclear markers, immunostaining was performed using the IntraPrep permeabilization kit (Beckman Coulter, Fullerton, CA, http://www.beckmancoulter.com) according to the manufacturer’s instructions. Primary antibody concentration was 1 μg per 1 million cells for mouse anti-RPE65 (Abcam) and mouse anti-MITF (C5) (Thermo Fisher Scientific) and 0.035 μg per 1 million cells for rabbit anti-OCT4 (Abcam). Secondary antibodies were either goat anti-mouse conjugated to Alexa 488 (Invitrogen) or goat anti-rabbit conjugated to Alexa 647 (Invitrogen). A nonspecific, species-appropriate isotype control was included in all flow cytometry experiments, and stained cells were analyzed using an Accuri C6 flow cytometer (BD Biosciences, San Diego, CA, http://www.bdbiosciences.com). For each marker, analyses were performed on three biological repeats.

**Phagocytosis Assay**

The cells were incubated for 16 hours either at 4°C or 37°C in ambient air with 0.1 mg per cm² of pH-Rhodo-labeled bioparticles (Invitrogen) resuspended in CO₂-independent medium (Invitrogen) supplemented with 4 mM glutamine (Invitrogen) and 1× antibiotic-antimycotic (Invitrogen). Afterwards, the cells either were fixed and immunostained or were dissociated and analyzed by flow cytometry.

**VEGF Enzyme-Linked Immunosorbent Assay**

RPE cells derived from hPSCs were grown on Transwell membrane (Corning) coated with Matrigel or VN-PAS Synthema II SC (Corning). Cell culture supernatants from the hPSC-RPE cell apical and basal sides, that is, upper and lower compartments of the Transwell, respectively, were collected after 48 hours of cell culture and sent on dry ice to Antigen Targeting and Consulting Service (Worcester, MA, http://www.atcs-incorporated.com). VEGF-A measurements were done in duplicate.

**Transplantation of Carboxyfluorescein Diacetate Succinimidyl Ester-Labeled hPSC-RPE Cells**

hESC-RPE cells were cultured for 20 days after plating, until they formed a lightly pigmented monolayer suitable for transplantation [30]. The cells were then labeled with 25 μM carboxyfluorescein diacetate, succinimidyl ester (CFSE) (CellTrace CFSE cell proliferation kit; Invitrogen) for 15 minutes at 37°C. The next day, the hESC-RPE cells were resuspended in RPE medium.

NOD-scid mice (NOD.CB17-Prkdcsid/J, 8–10 weeks old; Jackson Laboratory, Bar Harbor, ME, http://www.jax.org) were anesthetized by i.p. injection of a mixture of ketamine and xylazine. After making a hole at the pars plana with a 30-gauge sterile needle (BD Biosciences), a micropipette glass needle (tip internal diameter of 25–30 μm) mounted on a Pico-Injector holder (PU-100: Harvard Apparatus, Holliston, MA, www.harvardapparatus.com) was inserted through the hole into the vitreous, stopping at the surface of the retina. One microliter of the hESC-RPE cell suspension (5 × 10⁶/μl) was delivered into the subretinal space of each eye. Fundus photographs of the retina were taken every other day with a Micron III imager (Phoenix Research Lab, Inc., Pleasanton, CA, www.phoenixreslabs.com).

All of the surgical and animal care procedures were performed in compliance with the guidelines for care and use of laboratory animals of Johns Hopkins University and the Association for Research in Vision and Ophthalmology statement for the use of animals in ophthalmic and visual research.

**Reverse Transcription-Polymerase Chain Reaction and Quantitative Real-Time Polymerase Chain Reaction**

Total RNAs were extracted (RNeasy Mini Kit) and treated with RNase-free DNase I (both from Qiagen). Extracted RNAs were reverse-transcribed (High Capacity cDNA kit; Applied Biosystems, Foster City, CA, http://www.appliedbiosystems.com), reverse transcription-polymerase chain reactions (RT-PCRs) were performed with PCR SuperMix (Invitrogen), and quantitative
real-time PCRs (qPCRs) were performed with EvaGreen qPCR Mastermix (Abm, Richmond, BC, Canada, http://www.abmgood.com). Quantitative PCR samples were run in triplicate, and expression levels were normalized using the geometric mean of three reference genes: FBXL12, SRP72, and CREBBP [40]. Gene-specific primers (supplemental online Tables 1, 2) were obtained from published sequences or designed using Beacon Designer (PREMIER Biosoft, Palo Alto, CA, www.premierbiosoft.com). RNA from M1, a primary culture of RPE derived from an adult donor eye, was a kind gift from Dr. Noriko Esumi (Johns Hopkins University).

**Statistical Analysis**
Analysis of variance (Tukey test) analysis was done with Prism 6.01 (GraphPad Software, Inc., San Diego, CA, http://www.graphpad.com).

**RESULTS**

**hPSCs Maintain Pluripotency and Normal Karyotype After Long-Term Culture by Clonal Propagation**
Based on a previous report that hPSCs can be successfully propagated through the use of blebbistatin [33], we first checked whether the hiPSC and hESC lines that we were using could also be cultured long term following this approach. When cultured by clonal propagation, hPSCs grew from single cells (Fig. 1A) to large colonies ready for passage within approximately a week (Fig. 1A, 1B; supplemental online Fig. 1A). These colonies displayed morphological features typical of pluripotent cells, such as a smooth surface and tightly packed cells with large nuclei (Fig. 1C; supplemental online Fig. 1B). (Data obtained with Matrigel are reported in the main figures of the article, whereas data obtained with VN-PAS are shown in the supplemental figures except when otherwise noted.)
After more than 20 passages (>3 months) by clonal propagation, expression of key markers of pluripotency such as POU class 5 homeobox 1 (POUSF1, also known as OCT4), Nanog homeobox (NANOG), sex-determining region Y (SRY) box 2 (SOX2), and the cell surface antigen SSEA-4 were all observed in a majority of hPSC colonies, as assessed by immunostaining (Fig. 1D; supplemental online Fig. 1C). Confirming these results, flow cytometry analysis indicated that >97% of cells in culture stained positive for OCT4, whereas >91% were positive for the cell surface antigen TRA-1-60 (Fig. 1E; supplemental online Fig. 1D; supplemental online Table 3). Spontaneous differentiation was therefore minimal and comprised between 3% and 9% of the total cell population, according to the pluripotency marker assessed. Finally, chromosome integrity was evaluated by G-banding karyotype: for both hPSC lines, the majority of the 20 metaphase spreads analyzed were found to have a normal 46, XX karyotype (Fig. 1F; supplemental online Fig. 1E). Taken together, these data confirm earlier results and show that hPSCs can be cultured for extended periods of time on Matrigel or VN-PAS by clonal propagation while maintaining a high level of pluripotent marker expression and a stable karyotype.

Differentiation of hPSC Monolayers Is Associated With Progressive Expression of RPE Markers

In order to induce differentiation of hPSCs, cells were plated at high density (20,000 cells per cm²) in TeSR1 and cultured for 8 days, during which time the cells developed into a monolayer (Fig. 2A). At that point, the pluripotency maintaining medium was exchanged for differentiation medium (DM; see Materials and Methods). Following this protocol, spontaneously differentiating monolayers of hPSCs reproducibly generated RPE cells after several weeks [22, 23, 26]. For the first weeks, the differentiating monolayer remained colorless, and pigmented colonies typically started to be visible 25–30 days after the switch to DM (d1) (supplemental online Fig. 2). In order to understand the kinetics of RPE differentiation following d1, we monitored the expression of several markers by qPCR on a weekly basis (Fig. 2B). Expression of the pluripotency marker OCT4 was rapidly downregulated and became undetectable by week 3. Expression of the eye field transcription factor paired box 6 (PAX6) was clearly evident by week 1, with some increased expression over the next few weeks. Microphthalmia-associated transcription factor (MITF), a transcription factor (TF) required for RPE differentiation [41], showed a similar pattern of expression. In contrast, orthodenticle homeobox 2 (OTX2), another TF involved in RPE differentiation [41], was already highly expressed in hPSCs as previously [42], became downregulated from week 1 to week 2, and then showed little additional change. We also checked mature RPE markers and found that bestrophin 1 (BEST1), retinal dehydro-binding protein 1 (RLBP1), and retinal pigment epithelium-specific protein of 65 kDa (RPE65) all showed early detectable levels of expression that generally increased with time in DM. More specifically, BEST1, which encodes a chloride channel-related protein, was upregulated approximately three-to eightfold from week 1 to week 3, and RLBP1 and RPE65, which encode proteins responsible for the visual cycle, were both upregulated approximately fivefold during the same time interval. Tyrosinase (TYR), which encodes the enzyme responsible for the conversion of tyrosine to the pigment melanin, was undetectable during the first 2 weeks of differentiation. It is noteworthy that its expression increased markedly after the third week of differentiation, which correlated with the appearance of the first pigmented colonies in our cultures. Our results demonstrated that during the first 5 weeks of RPE differentiation, hPSCs gradually upregulate key RPE markers while downregulating the expression of pluripotency maintaining TFs.

Approximately 50 days after switching to DM, numerous large clusters of pigmented cells were readily observed in our cultures (Fig. 2C; supplemental online Fig. 2C). In order to assess the extent of differentiation toward putative RPE, we sought to analyze the cell monolayers by flow cytometry for expression of RPE65. Because monolayers of differentiating cells are difficult to dissociate into single-cell suspensions required for flow cytometry procedures, we first incubated them for an extended period of time in DM supplemented with collagenase IV. Upon prolonged incubation with collagenase, the monolayers detached and formed many small clumps. These clumps were then easily dissociated to single cells by treatment with Accumax. Following this approach, we were able to reproducibly obtain monolocular solutions of differentiating cells suitable for flow cytometry analysis. We found that the fraction of RPE65+ cells did not significantly differ between hiPSCs and hESCs, being 16.1 ± 0.2% and 16.6 ± 2.5%, respectively (Fig. 2D, 2E).

Enrichment of Putative RPE From hPSCs Leads to Monolayers of Pigmented Cells With Polygonal Morphology

Since it was possible to generate single-cell suspensions of differentiating hPSCs consisting of approximately 16% with RPE characteristics, we decided to further culture these cells. After 50 days in DM, whole dishes of differentiating hPSCs were passaged a first time (P1) by successive treatment with collagenase IV and Accumax (Fig. 2A). The cells were then plated at high density (100,000 cells per cm²) and cultured in RPE medium (see Material and Methods). Two weeks after P1, we observed that the dishes contained many large colonies consisting of lightly pigmented cells with characteristic cobblestone morphology (Fig. 3A). These colonies were intermingled with nonpigmented cells with a fibroblast-like morphology. In order to further enrich the culture for putative RPE cells, we thereafter passaged the whole dish a second time (P2) and cultured the cells as before. One month after P2, the cells had grown into a monolayer of lightly pigmented cells with cobblestone morphology (not shown). Surprisingly, there were no longer detectable nonpigmented fibroblast-like cells like those observed at P1. For characterization, the putative RPE monolayers were then allowed to mature for an additional 20 days (Fig. 2A). At that point, 50 days after P2, RPE cells were strongly pigmented and had a polygonal morphology (Fig. 3B, 3C; supplemental online Fig. 3A, 3B). The monolayer had formed many domes (not shown), suggesting that the cells were functional and engaged in active fluid transport, as is known to occur in mature tight junction-bearing RPE cells [3]. As an alternative way to maintain the hPSC-RPE cells, we were able to serially passage them every 15–20 days after P2.

hPSC-RPE Cells Form Polarized Monolayers and Express Key RPE Genes and Proteins

We used immunohistochemistry to characterize the expression of key RPE proteins by the enriched hPSC-RPE cell monolayers.
We detected strong expression of MITF and OTX2 (Fig. 3D; supplemental online Fig. 3C). In addition, we observed strong membrane staining of BEST1 and the tight junction protein ZO-1 (Fig. 3D; supplemental online Fig. 3C), which showed basolateral and apical localization, respectively (Fig. 3G, 3H; supplemental online Fig. 3D, 3E). This polarized expression of ZO-1 and BEST1 is in accordance with patterns described previously in both hPSC-RPE cells [22, 23, 26] and fetal RPE cells [36, 43] cultured in vitro, as...
wells as in adult RPE in vivo [44, 45]. Finally, the visual cycle proteins RLBP1 and RPE65 showed clear expression in most hPSC-RPE cells (Fig. 3D; supplemental online Fig. 3C). Interestingly, BEST1 and RLBP1 had a more heterogeneous staining pattern, similar to what was observed in hPSC-RPE cells obtained by manual picking (Fig. 3E). We also confirmed by Western blot that these mature RPE markers were expressed at comparable levels in hPSCs obtained by serial passage and manual picking (Fig. 3F). Taken together, these results demonstrate that hPSC-RPE cells form polarized monolayers that express many important RPE proteins.

Next, we further analyzed hPSC-RPE gene expression by RT-PCR. In addition to MITF and OTX2, detected previously by immunostaining, we also examined the expression of genes playing a role in RPE function (Fig. 4A). We observed strong expression of transcripts for TYR and premelanosome protein (PMEL), both involved in melanogenesis. Related to retinoid recycling, RLBP1, CRALBP, and lecithin retinol acyltransferase (LRAT) transcripts

Figure 3. Morphology of hPSC-RPE cells and expression of RPE marker proteins in pigmented monolayers. (A): Bright-field photomicrograph of hPSC-RPE colonies intermingled with fibroblast-like cells after passage 1. (B): A six-well plate with hPSC-RPE cells after 50 days of maturation in RPE medium. (C): Bright-field photomicrograph of hPSC-RPE cells shown in (B). (D): Expression of key RPE markers by immunostaining. Scale bars = 30 μm. (E): Expression of BEST1 and RLBP1 in hPSC-RPE cells obtained by manual picking. Scale bars = 30 μm. (F): Western blot analysis of BEST1 and RLBP1 expression. A protein standard was used on each Western blot to determine the correct molecular weight of proteins. (G, H): z-stack confocal micrographs showing typical polarized expression of RPE proteins, with ZO-1 (green) demonstrating apical localization (G) and BEST1 (green) demonstrating basolateral localization (H). The nuclei were counterstained with Hoechst (blue). Scale bars = 10 μm. Abbreviations: GAPDH, glyceraldehyde-3-phosphate dehydrogenase; hPSC, human pluripotent stem cell; Man., manual picking; RPE, retinal pigment epithelium; S.P., serial passage.
were detected. In addition to BEST1, transcripts for another membrane-associated protein, the extracellular matrix metalloproteinase inducer (EMMPRIN) [46], were expressed. Transcripts for pigment epithelium-derived factor (PEDF), an antiangiogenic neurotrophin secreted apically by RPE cells [47], were detected at high levels in hPSC-RPE cells, as were transcripts for c-mer proto-oncoprotein tyrosine kinase (MERTK), which is associated with phagocytosis [48]. Overall, hPSC-RPE cells cultured on Matrigel or VN-PAS express all the expected RPE markers that we tested for.

The extracellular matrix (ECM) has been shown to play an important role in RPE differentiation and hPSC-RPE gene expression [49]. We therefore sought to understand whether the two ECMs used in our study had an influence on RPE marker expression, as determined by qPCR. We did not detect any significant difference (p > .05) in gene expression for hESC-RPE or hiPSC-RPE cells grown on VN-PAS compared with Matrigel (Fig. 4B). Next, we compared mRNA expression levels between hPSC-RPE cells obtained after manual picking or those obtained after serial passage (Fig. 4B). Interestingly, we found PAX6 to be significantly upregulated, approximately 3.5-fold in hiPSC-RPE cells obtained by serial passage versus manual picking. On the other hand, PAX6 in hESC-RPE cells obtained by serial passage was downregulated approximately threefold compared with hESC-RPE cells obtained by manual picking. Since PAX6 expression is observed during RPE development in vivo but turned down as RPE matures [50], this result could suggest that for hiPSCs, serial passage may lead to RPE in a less mature state compared with manual picking, whereas the opposite may be true for hESCs. The other RPE markers analyzed showed minimal differences, with the
only differences above twofold not being statistically significant. However, there was a trend toward lower expression of the mature marker BEST1 in hiPSC-RPE cells obtained after serial passage (p values of .438 and .075 for manual vs. serial passage on Matrigel or vs. serial passage on VN-PAS, respectively), whereas the trend was toward higher expression levels for hESC-RPE cells isolated after serial passage (p values of <10^-4 and .1 for manual vs. serial passage on Matrigel or vs. serial passage on VN-PAS, respectively). Finally, we compared hiPSC-RPE cells obtained after serial passage with cultured rRPE cells and M1, a primary line of adult RPE cells [51] (Fig. 4B). hiPSC-RPE cells had overall lower mRNA levels for hESC-RPE cells isolated after serial passage, whereas the opposite was true for TYR, a finding similar to results reported previously [22]. With the exception of PAX6 in hiPSC-RPE cells and MITF in rRPE cells, gene expression levels were otherwise comparable between hPSC-RPE and native RPE cells for the other markers analyzed.

Together, these findings indicate that culturing hPSC-RPE cells on Matrigel versus VN-PAS does not significantly impact their gene expression profile, at least for the key RPE markers assessed. Similarly, RPE purification by serial passage did not significantly influence hPSC-RPE mRNA levels compared with hESC-RPE cells maintained on VN-PAS, where an opposite effect was observed in hESCs and hiPSCs. Finally, we did note differences in transcript abundance between hPSC-RPE and rRPE or M1, consistent with previous reports that hESC-derived RPE cells are not exactly equivalent to native RPE at the transcriptional level [52].

hPSC-RPE Cells Form Functional Monolayers Capable of Phagocytosis and Polarized Secretion of VEGF

An essential function assumed by RPE cells in vivo is the phagocytosis of outer segments shed by photoreceptors [3]. In order to test whether hPSC-RPE cells were capable of such phagocytosis, they were first incubated in the presence of pH-Rhodo-labeled bioparticles, fixed, and stained for the tight junction protein ZO-1, which marks the apical side of RPE cells. It is only upon entering low pH phagosomes that pH-Rhodo-labeled bioparticles become fluorescent, thus providing a convenient system to observe particles specifically engulfed by cells. Apically localized pH-Rhodo-labeled bioparticles were extensively observed within hPSC-RPE layers (Fig. 5A; supplemental online Fig. 4), indicating that these cells were capable of phagocytosis.

RPE cells in vivo are also known to secrete growth factors, such as VEGF, in a polarized fashion [3]. Similarly, native RPE monolayers in vitro show preferential secretion of VEGF to the basal side [43]. We therefore grew hPSC-RPE cells on Transwells and collected culture medium from both the upper and lower reservoirs, as has been described previously [53]. We found that VEGF was preferentially secreted by the basal side of hPSC-RPE cells, with basal to apical ratios ranging from 1.8 to 3.3 (Fig. 4C). Intriguingly, VEGF secretion levels were higher for hESC-RPE cells cultured on VN-PAS, with concentrations as high as 2,500 and 1,255 pg/ml for the basal and apical sides, respectively. In contrast, VEGF levels for hESC-RPE cells maintained on Matrigel were comparable to those observed for hiPSC-RPE cells with both ECMs; they ranged from close to 500 pg/ml up to 700 pg/ml for the basal side and between 250 and 380 pg/ml for the apical side.

Taken together, these results demonstrate that hESC-RPE and hiPSC-RPE cells generated through our new protocol, whether grown on Matrigel or on VN-PAS, are functional, showing both phagocytic activity and polarized growth factor secretion.

hPSC-RPE Monolayers Obtained After Serial Passage Have the Same Degree of Cell Purity as hPSC-RPE Cells Obtained After Manual Isolation

Since our protocol to obtain hPSC-RPE monolayers did not feature any isolation step but was rather based on progressive enrichment by serial passage, it was important to assess the degree of cell purity. To this end, hPSC-RPE monolayers obtained through serial passage or by manual picking were immunostained for MITF or RPE65 and analyzed by flow cytometry (Fig. 6A). hPSC-RPE cells obtained by serial passage were found to be 96.9%–99.1% positive for MITF, values similar to that observed with hPSC-RPE cells obtained by manual picking (99.3%–99.5%) (supplemental online Table 4). For RPE65, 97.6%–98.5% of cells expressed this marker in the case of hPSC-RPE cells cultured on Matrigel after both serial passage and manual picking. Interestingly, although hESC-RPE cells maintained on VN-PAS were over 98% positive for RPE65 expression, hiPSC-RPE cells maintained on the same surface demonstrated RPE65 expression in only 68% of the cells (supplemental online Fig. 4B). Surprised by this result, we repeated the experiment with hiPSC-RPE cells from a different batch of differentiated cells and obtained a similar fraction of RPE65+ cells (data not shown). This could mean that, in the case of hiPSCs differentiated on VN-PAS, enrichment by serial
passage is not complete, and a significant number of MITF+/RPE65− cells still contaminate the RPE monolayer. Visual observation under the microscope did not reveal any extended area of non-pigmented cells in hiPSC-RPE cells differentiated on VN-PAS, as would be expected from a contamination of almost 30% by non-RPE cells. Nevertheless, contamination could be due to melanocytes, which are MITF+ pigmented cells [54] expressing RPE65 protein at very low level compared with the RPE [55]. To test this possibility, PCR analysis for the melanocyte transcription factor PAX3 was carried out with the hiPSC-RPE cells grown on VN-PAS. The assay failed to detect any PAX3 expression (data not shown). Thus, the diminished ratio of RPE65+ cells does not seem to have been caused by melanocyte contamination; it more probably is a consequence of a lower level of RPE65 protein expression that is below the level detectable by our flow cytometry assay. Supporting this hypothesis, the median fluorescence intensity of RPE65 stained cells was approximately two times lower for hiPSC-RPE cells grown on VN-PAS versus Matrigel (supplemental online Fig. 4B).

To further characterize the functionality and consistency of the hPSC-RPE cells, we used a previously described phagocytosis assay [30]: pH-Rhodo-labeled bioparticles were seeded on hPSC-RPE monolayers, and the cells were incubated at 37°C and then analyzed by flow cytometry (Fig. 6B). Compared with controls incubated at 4°C, a temperature that inhibits phagocytosis, 89 and 83% of hESC-RPE cells cultured on Matrigel or VN-PAS, respectively, had phagocytosed pH-Rhodo-labeled bioparticles (Fig. 6C). This result was equivalent to that of hESC-RPE cells isolated by manual picking (87%) and close to that published in a previous study [30]. In contrast, the percentage of cells with phagocytosed particles was almost two times lower for hiPSC-RPE cells, ranging from 42% to 48%, regardless of the extracellular matrix used or the purification protocol followed. In summary, these data show that hPSC-RPE cells obtained after serial passage have a high degree of cell purity and that they are comparable to hPSC-RPE cells isolated by manual picking in terms of the fraction of functional RPE cells.

hPSC-RPE Cells Transplanted In Vivo Survive and Remain Functional

In order to test the ability of hPSC-RPE cells to survive in an in vivo setting, CFSE-labeled hESC-RPE cells were injected into the subretinal space of albino, NOD-scid mice. Fundus images taken 1 week later displayed the presence of numerous pigmented clusters in the injected area, suggesting that the hESC-RPE cells had survived the transplantation process (Fig. 7A). Retinal sections showed that hESC-RPE cells formed boluses in the subretinal space (Fig. 7B), as has been described previously at this time point [23]. Importantly, the presence of rhodopsin-positive fragments within the cellular contours of hESC-RPE cells (delineated by F-actin staining) indicates that these cells can phagocytose photoreceptor outer segments from the mouse retina (Fig. 7B, arrowheads).

DISCUSSION

We have described here, as a modification of a standard protocol for the isolation of RPE from spontaneously differentiating hPSC monolayers [18, 37], a method by which hPSCs are amplified by clonal propagation and in which RPE cells are enriched by serial passage instead of by mechanical picking. These modifications eliminate the need for the time- and labor-consuming manual steps usually required to culture hPSCs and to purify the RPE population and thereby provide a readily scalable approach to generate large...
numbers of high-quality RPE cells. A single six-well plate containing $1.2 \times 10^6$ hiPSCs, for instance, can produce up to $6 \times 10^7$ RPE cells at P2, which is up to 36 times more than the best protocols previously reported during the same time interval [22, 28, 30].

Since differentiation is achieved in monolayer, a standardized cell culture surface suitable for all stages of this process can be used to propagate hPSCs, obtain the initial RPE colonies, and further expand them. In addition, using simple culture media for RPE differentiation and culture, without the addition of serum, growth factors, or inhibitors, minimizes experimental variability while ensuring cost-efficient generation of differentiated cells. To our surprise, given the simplicity of our approach, we were able to obtain approximately 15% putative RPE cells after 50 days in differentiation medium, as assessed by RPE65 expression. RPE cell yield was not significantly different between our hiPSC and hESC lines, in accordance with previous studies [19, 52], whereas others reported lower efficiency with hiPSC lines [56, 57]. In the original report from Klimanskaya et al. [18], only 1% of EBs contained pigment cells after 4–8 weeks. Using different cocktails of growth factors or small molecules, several subsequent studies reported RPE yields ranging from 25% to 33%, after several weeks, as assessed by MITF expression or the presence of pigment cells [19, 21, 27, 58]. To date, the highest yield of RPE65+ cells observed after stepwise treatment with Noggin, retinoic acid, and Sonic Hedgehog was approximately 40% [59]. Thus, although the RPE differentiation efficiency following our protocol is lower than that reported in recent studies, it is not considerably so.

Obtaining a pure and functional population devoid of contaminating cells is a key requirement for the use of hPSC-derived RPE cells in biomedical applications [31]. Concerning hESC-RPE cells, analysis by flow cytometry showed that the RPE cells obtained after serial passage had a purity of 98%–99%, depending on the RPE marker chosen, which is essentially the same as that obtained by manual picking. The high degree of cell purity obtained with hESC-RPE cells cultured on Matrigel or VN-PAS is comparable to that reported in a previous study, where hESC-RPE cells were purified by mechanical dissection and injected into human patients enrolled in a clinical trial [30].

In vivo applications, such as regenerative therapy, would greatly benefit from more defined and xeno-free conditions for hPSC-RPE cell production, since it could limit unwanted animal byproduct-induced immunogenicity or contamination from animal pathogens [60]. Most published protocols include feeder cells for hPSC propagation or differentiation and sometimes fetal bovine serum [32]. Recently, two studies described defined [57]
or xeno-free [61] conditions for RPE differentiation, yet in both cases, hPSCs were maintained on a layer of feeder cells.

With the use of VN-PAS, completely defined conditions were achieved for both hESCs and RPE generation. Because hPSCs may be used to develop patient-specific therapy, we thought that it would be clinically relevant to reproduce with hPSCs the results obtained with hESCs on VN-PAS. To our surprise, although analysis by flow cytometry showed the percentage of cells positive for marker expression to be >96% for both MITF and RPE65 when hPSC-RPE cells were grown on Matrigel and for MITF when the cells were grown on VN-PAS, the percentage of RPE65+ cells cultured on VN-PAS was only 68%. Given the homogenous morphology of the hPSC-RPE culture on VN-PAS and the absence of PAX3 expression, contamination by melanocytes seems unlikely. Although we cannot formally exclude contamination by other MITF expressing cell types, an alternative hypothesis suggests that RPE cells from this specific hPSC line (hPSC IMR904) may not be maturing to the same level or at the same speed when grown on VN-PAS, compared with Matrigel. Further studies with an extended repertoire of hPSC lines will be required to determine whether this result is a more general property of hPSCs when VN-PAS is used for RPE generation.

In addition to marker expression, we also determined the fraction of functional cells within a given hPSC-RPE population. To this end we used a simple pH-Rhodo-labeled bioparticle phagocytosis assay described by Schwartz et al. [30]. With this approach, we observed similar percentages of functional hESC-RPE cells in their study and ours. By contrast, approximately 40% fewer hPSC-RPE cells had phagocytosed bioparticles: importantly, hPSC-RPE cells obtained by manual picking or serial passage performed equally, ruling out a possible negative effect from the latter procedure. These results are in contrast with a recent study of hPSC-RPE cells from the same line that showed that 80%–90% of them had internalized fluorescent latex beads [53]. In another report, no difference was found between hESC-RPE and hPSC-RPE cells for rod outer segment phagocytosis [22]. In both studies, different culture conditions were used for hPSC-RPE culture. Although it is known that growth medium [62] and extracellular matrixes [63] can have an influence on RPE phagocytic activity, it remains unclear why in our hands this muted effect is only observed with hPSC-RPE cells and not hESC-RPE cells.

Some variability was observed for PAX6 expression following RPE purification by serial passage, with a higher level in hPSC-RPE cells and a lower level in hESC-RPE cells, compared with RPE isolation by manual picking. Importantly, other key RPE transcription factors, such as MITF and OTX2, were left unchanged. Variable levels of PAX6 expression have been observed in hPSC-RPE cells to date: it was clearly detected with hPSC IMR90 and hESC H7 lines [18, 21–23, 27, 53], and in another study it was present with hESC lines but absent with hPSC lines [52]. These different hPSC-RPE cell lines nevertheless expressed mature RPE markers and were functional [18, 21–23, 27, 52, 53]. Similarly, despite different PAX6 expression levels, hESC-RPE and hPSC-RPE cells obtained by serial passage secreted growth factors in a similar fashion. They also displayed the same degree of phagocytosis as their counterparts obtained by manual picking. Therefore, the variation in PAX6 expression observed between hESC-RPE and hPSC-RPE cells after serial passage may not be important enough to affect their identity as RPE cells.

Comparison studies have suggested that hPSC-RPE cells display early senescence [22, 56] and differ more than hESC-RPE cells when compared with fetal RPE at the transcriptomic level [52]. However, others have demonstrated that hPSC-RPE cells display many RPE functions [31]. Thus, it would be of interest to investigate with a large sample of hESC and hPSC lines how RPE characteristics are affected when these cells are grown under a variety of conditions, so that differences inherent to ESCs and iPSCs may be uncovered. Such studies will be particularly relevant if hPSC-RPE cells were used as an in vitro model for drug screening or disease modeling.

**CONCLUSION**

In this report, we describe a new protocol for the generation of RPE from hPSCs, in which RPE cells are purified through two rounds of whole-dish single-cell dissociation. Together with clonal propagation of hPSCs, this differentiation process can be easily scaled up to allow the production of large amounts of highly purified and functional RPE cells. The monolayers of RPE cells demonstrated many characteristics of native RPE, including pigment and characteristic morphology, marker expression, polarized membrane and VEGF secretion, and phagocytic activity. We also show that the differentiation procedure can be achieved using totally defined conditions.

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**AUTHOR CONTRIBUTIONS**

J.M.: conception and design, collection and assembly of data, data analysis and interpretation, manuscript writing, final approval of manuscript; K.W.: collection and assembly of data, data analysis and interpretation, manuscript reviewing; D.G.: collection and assembly of data, data analysis and interpretation, manuscript writing, final approval of manuscript.

**DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST**

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