Corneal Spheres derived from Human Embryonic and Human Pluripotent Parthenogenetic Stem Cells

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Abstract
Corneal blindness is common. Cornea transplants are the most commonly performed organ transplants, but the need for corneal grafts worldwide far outweighs the supply of healthy donor corneas. Here we describe a differentiation protocol that yields corneal orbs from human embryonic stem cells (hESC) as well as from human pluripotent parthenogenetic stem cells (hpSC), and therefore can be manufactured free of transmissible pathogens. Cornea and other tissues generated from parthenogenetic stem cells that are homozygous at HLA loci have a distinct immunologic advantage over fully allogeneic grafts, and this report is the first to describe multilayered cornea generated from hpSC. The differentiated corneal product is layered and anatomically similar to normal human cornea, expresses appropriate corneal markers at the mRNA and protein (and secreted protein) levels, and is permeable to topical ophthalmic drugs. This 3D stem cell-derived cornea is a foundational step in development of appropriately organized, functional corneal grafts from hESC and hpSC for use in invitro assays as well as regenerative therapies.

Introduction
The cornea is more than a protective shield, it also serves as an external lens and is therefore critical for refraction and normal vision. An estimated 8 to 10 million worldwide are blind from corneal disease [1]. In developed countries, corneas are the most common ‘organ’ transplants, but a significant unmet medical need for corneal grafts persists. Fully artificial corneas have helped significant numbers of patients with severe corneal disease or injury, but do not meet the medical needs of most patients requiring corneal grafting, and so have limited impact on the need for donor corneas [2]. Stem cell-derived corneas, free of transmissible pathogens, represent a promising and safe alternative to cadaveric grafts, provided they can be generated cost-effectively, with function (including barrier, absorption, refraction) equivalent to that afforded by cadaveric grafts. Here we describe a novel multilayer corneal structure generated from human pluripotent stem cells, a first step in the goal of developing a full-thickness corneal graft from a single differentiation protocol.

The cornea is the clear (transparent) front part of the eye and is comprised of three main cellular layers - an outer non-keratinized stratified epithelium, middle stromal layer containing a hydrated extracellular matrix (ECM) with scattered corneal fibroblasts (keratocytes), and an inner, single-cell endothelial layer. Each layer is separated by distinct, specialized basal lamina. Bowman’s layer is most anterior, located between the epithelium and the stroma. Descemet’s membrane separates the stroma and the endothelium, serving as a basement membrane for corneal endothelium [3]. Advances in cornea engineering techniques have just started as another important alternative to overcome the limitation of 3D-corneal tissue available for transplantation [7,29]. To this end, bioengineered corneas to replace partial or full-thickness corneal defects have recently been developed as an alternative to cadaveric grafts [6,7] and an acellular collagen-based cornea was recently tested in a Phase I trial [8]. Decellularization of animal corneas is another promising method for the development of artificial human corneas by tissue engineering [37].

Adult human cornea harbors resident stem cells in the limbus (between the cornea and conjunctiva) [reviewed in 4], and these limbal stem cells mediate regeneration and wound repair of the corneal epithelium throughout adult life. Recent success in transplanting (autologous, adult) limbal stem cells for corneal burns [5] points to the importance of stem cell biology for therapy of corneal disease. Limbal stem cell transplantation, both autologous and allogeneic, is a major stem cell success story and is now available in many centers, but is not useful for diseases where more than corneal epithelium is compromised; complete 3D-corneas containing all cellular compartments of the cornea are still a necessity for treatment of the many patients with corneal blindness.

Derivation of human pluripotent embryonic stem cell (hESC) lines and recent advances in hESC biology have generated great interest in the field of stem cell-based cornea engineering [9-11]. Cell culture technologies that expand undifferentiated hESCs and subsequent methods to direct differentiation into corneal-like cells are improving and open up the possibility of producing corneal constructs for use in cellular therapy. Corneal epithelial cells have been produced by culturing hESC on collagen IV using medium conditioned by limbal fibroblasts[12].

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Our focus here is on differentiation of a whole cornea from pluripotent stem cells generally, but particularly human parthenogenetic stem cells or hpSC. hpSC are morphologically similar to hESC, express the same pluripotency markers, have high levels of alkaline phosphatase and telomerase activity, and give rise in vitro and in vivo to derivatives of all three embryonic germ layers. Unlike hESC, hpSC cannot differentiate into extraembryonic tissues [38,39]. Differentiation techniques during isolation of hpSC from unfertilized eggs allow creation of either heterozygous hpSC that are genetically identical to the oocyte donor, or HLA homozygous hpSC which are immunologically less complex [13-15]. As such, derivatives of these homozygous lines transplanted in appropriate donors will be less immunogenic than traditional cadaveric allografts. This report represents a first but significant step toward generation of sterile, functional, less immunogenic 3D corneal grafts for use in regenerative therapies.

Materials and Methods

In vitro differentiation of corneas from human pluripotent stem cells

The WA09 hESC line was provided by Dr. Mike West. The hpSC lines hpSC-1, hpSC-3, hpSC-5 [13] and hpSC-Hhom-1 [14] were derived by Revazova et al., and are owned by International Stem Cell Corporation. Undifferentiated stem cells were maintained on a feeder layer of human Mitomycin-C treated dermal fibroblasts (MCFibs”; Lifeline Cell Technology), Frederick, MD) in hESC Growth Medium (Knockout™ D-MEM; Invitrogen, Carlsbad, CA) supplemented with 2.4 mM GlutaMax (Invitrogen), 0.12 mM MEM Non-Essential Amino Acid Solution (Invitrogen), 0.12% mercaptoethanol (Invitrogen), 8% Plasmanate, 8% Knockout Serum Replacement (Invitrogen) 10 ng/ml basic fibroblast growth factor (bFGF, Lifeline Cell Technology), and 10 ng/ml leukemia inhibitory factor (LIF, Chemicon, Billerica MA).

To initiate differentiation the stem cells were allowed to grow for approximately 10 days (with medium replacement every other day) in a humidified 5%CO₂ incubator. When colonies formed (morphologically similar to burst forming units with sharp edges), both LIF and bFGF were removed from the medium, and on day 17, these differentiated cultures were passaged with Collagenase Type IV (890 U/ml; Lifeline Cell Technology) and transferred to deep 6-well plates (Greiner Bio-One, Frickenhausen, Germany) for further differentiation in low-attachment conditions. The cultures were maintained for an additional 100-120 days in a humidified atmosphere of 5% CO₂/95% air, with medium added twice a week, during which the dome-like colonies grew in size and became fluid-filled clear spheres. The spheres break off from the adherent cells and float freely; these were retrieved from the medium for characterization.

Histology and immunocytochemistry

Ten corneal orb constructs derived from hESC or hpSC were used for this study. For histological analysis the corneal orbs were cut in half, each half measuring approximately 5 x 10 mm. One half was fixed for 12 hours in 10% buffered formalin, embedded in paraffin and sectioned to a thickness of 5 μm (small size hpSC-derived orbs were fixed in their entirety). Sections were stained with hematoxylin and eosin (H&E), Masson’s trichrome, or periodic acid Schiff (PAS) using standard staining protocols. Sections were also stained for cytokeratins, using primary AE1/3 antibody (Chemicon) for pan-cytokeratins.

For immunofluorescence assays, orb halves (or whole small size constructs) were snap-frozen in OCT compound and sectioned on a Tissue Tek II Cryostat. Frozen sections (5 μm) were air-dried and fixed with cold (-20°C) acetone for 10 minutes, then blocked with 1% bovine serum albumin (BSA) in PBS at room temperature for 1 hour. Sections were incubated with primary antibodies at 4°C overnight and subsequently with appropriate secondary antibodies at room temperature for 45 minutes. Primary antibodies were: mouse anti-keratin 3, 1:50 dilution (AE3 clone, MP Biomedicales, Solon OH), mouse anti-cytokeratin 18, 1:50 (Santa Cruz Biotechnology Inc., Santa Cruz CA), mouse anti-cytokeratin 19, 1:200 (Abcam, Cambridge MA), rabbit anti-connexin-43, 1:100 (Invitrogen), rabbit anti-mucin-1, 1:100 (Abcam), rabbit anti-zona occludens (ZO-1), 1:50 (Invitrogen). Secondary donkey anti-mouse or anti-rabbit antibodies conjugated with AlexaFluor®546 (Invitrogen) were used. For vimentin staining, fixed tissue sections were embedded in paraffin and microwaved for antigen retrieval. Slides were washed briefly in PBS and then after blocking and permeabilization, treated with primary monoclonal antibody (Clone LN-6, Sigma-Aldrich, St. Louis MO). After labeling with the appropriate secondary antibodies, and nuclear counterstain with 4',6-diamidino-2-phenylindole (DAPI), or propidium iodide, all sections were examined using fluorescence microscopy. Positive controls for these studies were cadaveric human corneas.

Preparation of human corneal samples

Non-diseased banked human corneas (n = 2) were obtained from San Diego Eye Bank. The eyes were procured and processed using criteria established by the Medical Standards of the Eye Bank Association of America and the U.S. Food & Drug Administration.

Real-time quantitative PCR (RT-qPCR)

Stem cell-derived corneal constructs were examined for expression of normal human corneal genes using real-time quantitative PCR (RT-qPCR). Total RNA was extracted using the QiAasympomatic purification system, according to the manufacturer’s instructions (Qiagen, Valencia CA), and 100-500 ng total RNA was used for reverse transcription with the iScript™ cDNA synthesis kit (Bio-Rad,). PCR reactions were run in duplicate using 1/40-th of the cDNA per reaction and 400 nM forward and reverse primers or the Quant iTect™ Primer Assay, together with Quant iTest SYBR® Green master mix (Qiagen). Real-time PCR reactions were run on the Rotor-Gene® Q (Qiagen). Relative quantification was performed against a standard curve and normalized to the expression levels of one of the following housekeeping genes: cyclin G (CYCG), beta-glucuronidase (GUSB) or TATA box binding protein (TBP). After normalization, the samples were plotted relative to the first sample in the data set and the standard deviation of the expression measurements was calculated. The source of primers is shown are shown in Table 1. RNA isolated from human donor cornea for use as a comparison for gene expression of stem cell-derived corneas.

Drug permeability of stem cell-derived corneas

Corneal disease is often treated with topical drugs, possible because the normal cornea is permeable and allows rapid absorption [reviewed in 16]. The goal of these first permeability studies was qualitative, to test whether compounds with known high and low absorption behaved as expected relative to each other when applied to the differentiated corneal orbs. To test permeability function, intact corneal orbs were incubated with test compounds (10 μM) at 37°C for 30 min. Two model compounds were used – atenolol and antipyrine, as low- and high-permeability reference compounds, respectively. After incubation, the
fluid inside the orbs was carefully collected by needle aspiration, and used to measure drug concentrations by LC-MS/MS. The apparent permeability coefficient ($P_{\text{app}}$), a parameter commonly used to express in vitro permeability across a cell monolayer or tissue, was calculated as: $P_{\text{app}} = \frac{Q}{C_{\text{in}} \times A \times T}$, where $Q$ is the amount of drug accumulated in the orbs, $C_{\text{in}}$ is the applied drug concentration, $A$ is the surface area of the orbs, and $T$ is the incubation time.

**Determination of growth factor levels in differentiating cornea conditioned medium**

Commercial TGF-β1 and EGF ELISA immunoassay kits (Invitrogen) were used for quantitation of these factors in conditioned medium from cultured corneal constructs at different stages of differentiation. The conditioned medium was diluted 1:1 with serum-free DMEM, and assayed according to kit manufacturer’s protocols.

**Results**

**Differentiation and morphology of stem cell-derived corneal orbs**

The typical early appearance of cells that differentiated into corneal orbs was as small, tightly packed colonies in which individual cells were round, in comparison to the spindle-shaped feeder layer cells (Figure 1A). Often a few healthy stem cell colonies were noticeable the first day after low density inoculation, and became more heterogenous in shape, containing flattened cells with epithelial morphology, attached to the culture dishes. (Figure 1B,C). After 5-7 days more of differentiation, the colonies became multilayered, dense clusters in which individual cells were indistinguishable by light microscopy. In some colonies, a distinct clear spherical (orb-shaped) dome was observed, indicating the formation of a more mature cornea-like structure (Figure 1D). On day 17-19 the colonies were subcultured onto deep well plates under low attachment conditions, and within the next 3-4 weeks, freely floating pellucid spheres (corneal orbs) could be observed under a dissecting microscope.

**Figure 1**: Corneal orb differentiation from pluripotent human stem cells.

A. Early appearance of pluripotent stem cell colonies that will go on to form corneal orbs. The small, tightly packed cells in the colony are distinguishable morphologically from the spindle-shaped feeder cells. Scale bar = 300 µm.

B. Proliferation of the pre-cornea colonies results in larger colonies of irregular shapes but with clear edges demarcating the colony from spindle-shaped feeder layer cells. Scale bar = 300 µm.

C. Another expanding colony of differentiating cells easily distinguishable by light microscopy from the feeder layer. Scale bar = 300 µm.

D. Higher power appearance of differentiated pluripotent cells just before the cells are passaged onto low-adherence, deep plates. Scale bar = 100 µm.

E. Differentiating hpSC from which small (~1-2 mm) orbs have emerged. Scale bar = 1500 µm.

F. Mature corneal orb after 120 days of differentiation is translucent and fluid-filled.

G. Large, mature corneal orb after 120 days of differentiation with diameter of ~15mm.

Table 1:

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Histological analysis demonstrated that the spheres have interior and outer layers similar in arrangement to normal human cornea. These first histological studies indicated that the epithelium of corneal constructs began as a single layer (Figure 2A) and progressively developed additional layers (Figure 1B), following normal developmental patterns. This epithelial layer was PAS-positive indicating glycogen or other polysaccharide content (Figure 2C). PAS stains are used in pathologic evaluation of corneas to visualize corneal epithelial basement membrane (and Descemet’s membrane). Histologically the largest layer of the corneal orbs was the stromal layer, morphologically consistent with fibrous tissue. Masson’s trichrome staining (Figure 2D) shows that the stromal layer contains collagen fibrils (blue) that were heterogeneous in diameter, some disorganized fibrin strands (pink), and the subepithelial fibrotic membrane, which stained red. (Retention of the red in trichome staining may indicate a poorly permeable structure.) Cytokeratin staining (Figure 2E) was used to further analyze the corneal orbs (using a pan-cytokeratin AE1/3 antibody). Flattened (pantcytokeratin-negative) cells were found dispersed between the collagen fibrils, consistent with corneal fibroblasts or keratocytes. The most intense staining was in the superficial epithelial layer (seen at higher power in Figure 2E,F). In addition, in some orbs we identified a thin membrane layer (PAS-negative), between the outer layer of epithelium and the large stromal layer, suggestive of a developing Bowman’s membrane but this membrane was not fully formed and was not present in all orbs. The innermost surface of the orbs, where endothelium should be, was a single-cell layer with no evidence of
keratinization (by pan-cytokeratin staining). Routine hematoxylin and eosin (H&E) staining was consistent with an endothelium but this stain is non-specific.

Overall, histologic evaluation of these novel stem cell-derived orbs indicated significant structural similarity to human cornea, though a distinct endothelium could not be identified. In addition cultivation of the orbs at an air-liquid interface resulted in thickening of the epithelial layer (comparing Figures 2E and 2F), similar to increased stratification of human corneal epithelium during development at the time of eyelid opening [17,18].

Immunocytochemistry of cornea-like constructs confirmed expression of typical cornea proteins, and normal adult corneas were used to validate the staining pattern. Stem cell-derived corneas expressed mucin-1, a marker of corneal stratified epithelium (and mucin is a component of tears), throughout the epithelial layer (Figure 3A, normal corneal 3B), and the gap junction protein, connexin-43 (Figure 3C, normal cornea 3D), most strongly by the suprabasal (central) epithelial cells. Cytokeratins (using a pancytokeratin antibody, Figure 3E and normal cornea 3D), were also expressed, as expected. Specific cytokeratin expression was also examined, and the orbs expressed cytokeratin 19 at low levels (Figure 1G and normal cornea 1H) and cytokeratin 18 (Figure 1I) for which staining was intense only in the superficial layers of the corneal orbs. Strongly positive staining for the tight junction protein ZO-1 was found in all studied specimens, with appropriate subcellular localization at cell boundaries suggesting the formation of tight junction complexes, a feature of endothelial cell differentiation (Figure 1J). ZO-1 and connexin-43 staining patterns in the corneal orbs and normal human corneas were very similar, suggesting maturation of intercellular junctions in stem cell-derived orbs. Vimentin staining was positioned in all orbs at low levels, restricted to some stromal cells and the cell layer at the normal anatomic site (innermost) of corneal endothelium (Figure 1K). Stratified epithelial cells were vimentin-negative.

Gene expression of corneal markers

RT-qPCR assays were used to compare corneal gene expression in fully developed human cornea vs. stem cell-derived corneal orbs. Several genes known to function in all three major layers of the cornea were chosen for study (Figure 4). The expressions of collagen 5 (Col5), decorin, biglycan, lumican, keratocan, CK8, and CK18 in all samples analyzed were significantly higher in stem-cell derived corneas than in adult cornea. The corneal orbs also expressed many characteristic markers of human corneal epithelium including importin 13, integrin alpha 9, connexin-43, and enolase-a. Increased expression (vs. adult cornea) of ZO-1, confirming impressions from immunocytochemistry, was also noted in all stem cell-derived corneal orbs.

Drug permeability of corneal orbs

Permeability and transcorneal transport of drugs are critical processes in pharmacologic treatment of eye disease. To evaluate these functions we performed a small pilot study with two drugs normally delivered topically to the eye. From these studies the permeability coefficient (Papp) of the beta-blocker atenolol was calculated to be 2.60 + 1.55 (mean ± standard deviation) and the Papp of antipyrine was 15.5 + 13.1. Thus, the low vs. high permeability drugs were transported as expected relative to each other by the stem-cell derived corneas. Determination of growth factors in differentiating cornea conditioned medium (ELISA)

Corneal-derived EGF is long known as an important factor in corneal epithelial regeneration [reviewed in 19], while TGFβ1 has pleiotropic effects in the cornea including maintenance of the stromal layer [20]. Before ninety days of differentiation, neither EGF (in 4/4 samples) nor TGFβ1 (in 2/2 samples) was detectable using ELISA assays. After that time point, 3 of 4 samples had detectable EGF; the concentrations were 4.3, 0.8, and 2.2 pg/mL (at the lower limits of detection of the assay). TGFβ1 was detectable in 3 of 5 samples after 90 days of differentiation; the concentrations were 9.8, 20.6, and 2.8 pg/mL (at the lower limits of detection of the assay).

Discussion

The 3D corneal constructs generated in this work emerge as spheres from pluripotent stem cells with several important features of normal cornea including the basic anatomic layering, gene and protein expression patterns, rapid permeability to ophthalmic drugs, and no obvious opacity. This 3D structure represents a significant advance, since generation of functional miniorgans from pluripotent stem cells is very unusual. When induced to differentiate without engineering scaffolds or support, differentiation usually produces random collections of cells in culture, without functional organization. In the case of cornea, the anatomic organization is especially notable, since during normal corneal development, the stroma is derived from neural crest (migrating in to periocular mesenchyme [Hay 1979] whereas local ectoderm (adjacent to lens) becomes corneal epithelium (Cintron,
Figure 3: Fluorescent micrographs of immunostaining performed on corneal orbs at the end of differentiation process. DAPI in A-J marks nuclei in blue, propidium iodide in red (K). All scale bars are 300 µm.

A. Mucin-1 immunostaining (red) of corneal orb concentrated in the epithelial layer is stronger

B. Mucin-1 immunostaining (red) of normal adult human donor cornea.

C. Connexin-43 immunostaining (red) of corneal orb is concentrated on the innermost layer of epithelial cells and overall staining is more intense than

D. Connexin-43 staining (red) of normal adult human cornea.

E. Pancytokeratin immunostaining (red) is intense in the epithelial layer of differentiating corneal orbs (confirming patterns in Figure 2).

F. Less intense and more diffuse pancytokeratin staining (red) in normal human adult cornea than in differentiating orbs.

G. Cytokeratin 19 immunostaining (red) of corneal orbs is intense in the most superficial and deepest epithelial cell layers of the differentiating orbs.

H. Cytokeratin 19 immunostaining (red) is most intense in the most superficial epithelial cell layer of normal human adult cornea.

I. Cytokeratin 18 (red) is abundant in the most superficial epithelial cell layer of differentiating corneal orbs.

J. ZO-1 staining (red) is intense in differentiating corneal orbs with expected subcellular localization to the cell-cell edge interfaces indicating the presence of endothelium.

K. Vimentin negative staining (red) of the multilayered epithelial cells (arrow 1) and the vimentin positive-staining (green) of differentiating corneal orbs is intense in the entire epithelial stromal layer (interspersed stromal keratocytes, arrow 2) and at the predicted anatomic location of the endothelial monolayer (arrow 3).
Figure 4: Trends in relative gene expression of corneal genes in differentiating corneal orbs (n=3 independent orbs) vs. adult human cornea (n=2 independent corneas) show that the differentiating orbs express many normal corneal genes. Data are expressed as mean and bars are standard deviation. (Statistics not performed because of low n and high S.D.)

A. Importin 13 may be expressed by a resident limbal stem cell population in both differentiating and normal adult corneas, but the limbal stem cell population was not assayed with other markers. Integrin alpha 9, connexin-43, and enolase-α are normally expressed in corneal epithelium.

B. Connexin-43 and ZO-1 are both important for cell-cell communication in the cornea, and combined with immunohistochemistry (Figure 3) these results suggest maturation of subcellular structures involved in intercellular communication by 120 days of differentiation.

C. Differential expression of cytokeratins suggests relatively high expression of CK8 and CK19 in the stem cell-derived differentiating corneal orbs relative to mature adult cornea.

D. The most striking pattern in the gene expression studies was the high expression of various extracellular matrix proteins in the differentiating orbs compared to the lower level expression in mature adult cornea. At the protein level these ECM genes are low abundance in normal cornea compared to the relatively large collagen content of cornea.
1983), yet both layers were generated in a single differentiation protocol from pluripotent stem cells. Using the conditions we describe, generation of the epithelial layer in the corneal orbs appears to be robust by histology, by functional manipulation of the epithelium at an air-liquid interface, supported by expression of multiple developing epithelial genes (cytokeratins 8, 10, and 18, mucin-1, connexin-43, and enolase-a). Generation of the large stromal layer is also consistent, as seen histologically, by PAS staining, and gene expression (decorin, lumican, biglycan, and keratocan). The strona appears to be quite differentiated and lacked expression of the early stromal marker, Pax6. Some optimization is likely required for the endothelial layer, which may require longer differentiation (or an added differentiation step) to develop. In some corneal orbs we identified cells histologically that were appropriately localized and shaped for endothelial cells. Vimentin and zona ocludens staining (expressed during endothelial development) also indicate some endothelial cells in the orb interior, but we cannot yet determine if endothelium is reliably incorporated into the corneas at 100-120 days of differentiation. We have not yet quantified the number of limbal stem cells in the differentiating corneas, but their presence is suggested by the expression of importin-13 (IPO13) characteristic of corneal epithelial progenitor cells. In normal cornea, IPO13 is uniquely expressed by human limbal basal epithelial cells, and plays an important role in maintaining the progenitor phenotype and high proliferative potential [21].

In a small pilot study we examined permeability of the corneal orbs. The primary pathway of drug permeation from the surface tear fluid to the anterior chamber of the eye is transcorneal [22] and passage through the corneal epithelium is considered to be the rate-limiting step in the transcorneal penetration of most ophthalmic drugs. Our results suggested rapid corneal permeability of the two drugs tested and as well as the expected greater permeability of antipyrine over atenolol. Some secretory function of the corneas was confirmed at the later stages of differentiation. EGF, which stimulates corneal epithelial cell proliferation and migration [reviewed in 23] and TGFβ1, involved in maintaining corneal integrity, in part by counterbalancing the stimulatory effects of EGF [reviewed in 24] were detectable in medium around the developing corneas. Both EGF and TGFβ1 promote keratocyte differentiation [25] and chemotaxis [26].

Though the generation of a 3D cornea from a single cell source rather than via multiple differentiation procedures is somewhat surprising and promising for clinical manufacture, the pluripotent stem cell-derived corneas will also require further manipulation to optimize them for preclinical studies, particularly to perfect the alignment of collagen in the stromal layer to assure optimal refractive properties [27]. Normal corneal fibroblasts produce collagen fibrils of uniform size and spacing, which are then orthogonally arranged in the stroma matrix [28], and changes to this pattern of organization result in loss of corneal transparency. Finally, though the corneal spheres appear to be transparent, their true transparency needs to be quantified by physical measures.

The differentiation process yields corneal orbs with sufficient integrity that we anticipate they can be manipulated in engineering studies and moved to other culture (or co-culture) environments for further differentiation. The orbs are also suitable, after some further development, for combination with various bioengineered corneal materials in a ‘combined device-cell therapy’ approach taken by other groups [6,29]. These are all research agendas we are currently investigating, to take advantage of recent progress in generating biosynthetic corneas. Among the most promising of these biosynthetic corneas, a recent Phase I trial of cross-linked collagen corneal grafts were implanted into ten patients (9 with keratoconus) who had intact corneal endothelium. The grafts induced endogenous re-epithelization over the course of about a month, re-innervation starting at about a year, supported normal tear formation, without prolonged immunosuppression and without pain. Most of the treated patients could tolerate contact lenses to improve vision, which was not as good as the vision of comparable patients who received full thickness human allogeneic grafts [8].

The smaller size (though similar organization) of corneal orbs generated from hpSC compared to those generated from hESC will require the protocol for hpSC undergo modification relative to the protocol used with hESC. We have reported other differences in growth and differentiation patterns between hpSC and hESC lines. For example, when a protocol that yields efficient (~80%) differentiation of hESC into definitive endoderm [30] is applied to hpSC, the efficiency is about half that of hESC. But pre-treatment of the hpSC with trichostatin A increases the efficiency (~75%) to that of hESC differentiation [31]. The yield of neural progenitor cells from hpSC is also less than from hESC, which may depend in part on differences in expression of extracellular matrix proteins necessary to hold neurospheres together during differentiation [32]. Adaptation of protocols that do not require neurospheres is therefore a rational choice for hpSC. Differentiation of retinal pigment epithelial (RPE) cells from hpSC and hESC, accomplished without an intermediate sphere formation step, yields similar numbers of qualitatively similar RPE [32].

The focus on parthenogenetic stem cells as a source for corneas (and other tissues), is well-justified by the clinical literature on the immunogenicity of allogeneic corneal grafts. Rejection of cornea allografts is a major cause of graft failure [33], and a history of rejection puts patients at high risk of future (second) graft loss [34]. HLA matching can improve corneal allograft survival [35] but is not routinely done, based in part on cost, and because there is not global consensus on its value. Nonetheless, the major cause of graft failure in the first year after corneal transplantation is cell-mediated rejection [36]. A bank of HLA homozygous parthenogenetic stem cells to cover common HLA haplotypes in the population is, therefore, a potentially valuable source of transplantable cells that can be better matched to reduce the intensity and incidence of rejection.

In summary, we generated free-floating corneal orbs from both human ESC and smaller orbs from parthenogenetic pluripotent stem cells, and these corneas differentiate as a 3D layered structure similar to that of normal human corneas. A priority for continued pre-clinical development of these corneas is engineering approaches to improve stromal collagen organization and on developing variances in culture protocols aimed at increasing the size of hpSC-generated orbs. The clinical need for corneal grafts for blinding diseases, especially outside the U.S., is great, highlighting the importance of further research to optimize these potential stem cell-derived grafts.

Acknowledgments

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References


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