Isolation and Expansion of Multipotent Progenitors from Human Trabecular Meshwork

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To expand multi-potent progenitors from human trabecular meshwork (TM), we have created a new optimized method on two-dimensional (2D) followed by three-dimensional (3D) Matrigel in modified embryonic stem cell medium supplemented with 5% fetal bovine serum (MESCM + 5% FBS). The expanded TM cells were small cuboidal cells expressing TM markers such as AQP1, MGP, CHI3L1, and AnkG, embryonic stem cell (ESC) markers such as Oct4, Sox2, Nanog, and ABCG2, and neural crest (NC) markers such as p75NTR, FOXD3, Sox9, Sox10, and MSX1. Although expanded cells lost expression of these markers after passage, the cells regained the markers when Passage 2 cells were seeded on 3D Matrigel through activation of canonical BMP signaling. Such restored progenitors could differentiate into corneal endothelial cells, adipocytes, and chondrocytes but not keratocytes or osteocytes. Therefore, we have concluded that human TM harbors multipotent progenitors that can be effectively isolated and expanded using 2D Matrigel in MESCM + 5% FBS. This unique in vitro model system can be used to understand how TM is altered in glaucoma and whether such TM progenitor cells might one day be used for treating glaucoma or corneal endothelial dysfunction.

The main outflow pathway for aqueous humor in the eye consists of a series of endothelial cell–lined channels in the angle of the anterior chamber comprising the trabecular meshwork (TM), Schlemm's canal, the collector channels, and the episcleral venous system. The TM, especially the juxtacanalicular region and the inner wall of Schlemm's canal are thought to be the source of the most resistance to aqueous outflow. Reduced cellularity and function within the TM is observed with age and correlates with increased outflow resistance and elevated intraocular pressure (IOP)1–3. Consequently, dysfunction of TM cells might play a role in blindness caused by glaucoma4.

As a first step to explore the pathogenic role of TM cells in glaucoma, it is necessary to isolate and expand TM cells in vitro. In this regard, cumulative evidence suggests that there is a population of progenitors in Schwalbe's Ring, which is the transition area between the periphery of the corneal endothelium and the anterior non-filtering portion of the TM5. This hypothesis primarily comes from the following observations: (1) an increase in TM cell division is localized to the anterior non-filtering portion of the TM after argon laser trabeculoplasty6 and (2) stem cell markers such as nestin, alkaline phosphatase and telomerase are expressed in cells in the transition region7. In addition, Du et al.8,9 have isolated human TM cells by collagenase digestion and cultured on plastic in stem cell growth medium (SCGM) and such cells expressed ABCG2, Notch1, Oct4 and MUC1. Primarily isolated and expanded side population of TM cells can then differentiate into keratocytes and expanded TM cells can home to mouse TM and differentiate into TM cells in vivo9, suggesting that these cells might be progenitor cells.

Herein, we report our effort in optimizing the method of expanding TM progenitor cells using 2D Matrigel and MESCM + 5% FBS. Our method yields small cuboidal TM progenitors expressing markers of TM, embryonic stem (ES), and NC cells. Unlike what has been reported by Du et al.8, these progenitor cells can be differentiated into corneal endothelial cells, adipocytes, and chondrocytes but not keratocytes or osteocytes. Interestingly, maintenance of such a progenitor status requires activation of the canonical BMP signaling. The significance of these findings in studying the pathogenesis and being deployed as a cell-based therapy for glaucoma and corneal endothelial dysfunction is further discussed below.

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Results

2D Matrigel is preferred to maintain TM morphology and phenotype. A number of substrates such as fibronectin, gelatin (collagen) and plastic without coating have been used to expand TM cells. A number of media have also been used to culture TM cells including Dulbecco’s modified Eagle’s Medium (DMEM) + 10% FBS, Medium 199E + 20% FBS, Improved Minimum Essential Medium (IMEM) + 10–20% FBS, and SCGM (a stem cell culture medium with 5% FBS). Because we have successfully expanded human limbal niche cells on 2D Matrigel in MESCM, we thus compared four substrates, i.e., plastic, fibronectin, collagen IV, and 2D Matrigel and three media, i.e., MESCM, MESCM + 5% FBS, and SCGM. When cultured in the same medium, i.e., MESCM + 5% FBS, freshly isolated TM cells turned to “spindle” fibroblastic-like cells after 4 weeks of culture on plastic, fibronectin, and collagen IV but remained cuboidal on 2D Matrigel (Fig. 1A, top panel). Under this culturing condition, the transcript expression of TM markers (such as AQP1, CHI3L1, MGP, and AnkG) was maintained when cultured on 2D Matrigel or fibronectin but significantly reduced on plastic or collagen IV (Fig. 1B), and the immunostaining data also showed the same result (Fig. 1F, top four panel). However, the transcript expression of ESC and neural crest markers by cells cultured on 2D Matrigel was significantly higher than those cultured on fibronectin, plastic, or collagen IV (Fig. 1D). These results suggest that 2D Matrigel was the preferred substrate to maintain TM morphology and phenotype when cultured in MESCM + 5% FBS.

When cells cultured on 2D Matrigel was changed from MESCM + 5% FBS to SCGM, they also turned into spindle cells (Fig. 1A, bottom panel). Cells remained somewhat cuboidal in MESCM without 5% FBS (Fig. 1A, middle panel) and had similar transcript expression of TM cell markers (such as AQP1, MGP and stromal marker Vim) but significantly higher transcription expression of CHI3L1 and AnkG (Fig. 1C), which was confirmed by immunostaining (Fig. 1F). However, the cell’s transcript expression of ESC and NC markers such as ABCG2, KLF4, Nanog, Nestin, Oct4, Rex1, Sox2, SSEA4, FOXD3, HNK1, MSX1, Sox10 and N-cadherin was significantly less than those cultured in MESCM + 5% FBS (Fig. 1E).

MESCM + 5% FBS is preferred to promote expansion. Although both MESCM and MESCM + 5% FBS were able to maintain the TM phenotype in freshly isolated TM cells (Fig. 1), TM cells cultured in MESCM without FBS did not grow well at P1 and stopped growing at P3 with an enlarged shape and cell debris (Fig. 2A, top panel, and Fig. S1), resulting in a total cell doubling of 7 (Fig. 2B). In contrast, TM cells cultured in MESCM + 5% FBS could be expanded to 8 passages with a total cell doubling of 16 (Fig. 2C). Hence, we concluded that expansion of TM cells was promoted in MESCM + 5% FBS, but not MESCM, when seeded on 2D Matrigel.

However, there was a gradual loss of the TM phenotype during serial passages on 2D Matrigel in MESCM + 5% FBS as evidenced by significant reduction of transcript expression of TM markers such as AQP1, CHI3L1, MGP, AnkG and the stromal marker Vim (Fig. 3A). Immunostaining showed that freshly isolated TM...
cells (D0) positively expressed all these four markers (Fig. 3B) similar to what have been previously reported\(^6,19\). However, TM cells exhibited reduced expression of MGP and AnkG more so than CHI3L1 and AQP1 after Passage 1 (Fig. 3B). In addition, the loss of the TM phenotype was also accompanied by significant reduction of...

Figure 2. MESCM + 5% FBS is preferred to promote expansion. TM cells were expanded on 2D Matrigel in MESCM or MESCM + 5% FBS and each passage was subjected to morphological analysis by phase-contrast microscopy (A, scale bar: 100 µm). Cells expanded in MESCM alone (B) yielded fewer passages and total cell doubling than those expanded in MESCM + 5% FBS (C, n = 3, *P < 0.05 when compared to that of P0 cells).

Figure 3. Expanded TM cells lose their phenotype and progenitor status on 2D Matrigel. Freshly isolated TM cells (D0) and TM cells cultured on 2D Matrigel in MESCM + 5% FBS were subjected to qRT-PCR for transcription expression of TM markers (A) and ESC and neural crest markers (C, using freshly isolated TM cells as the control, n = 3, *P < 0.05) and immunostaining of TM markers (B, scale bars: 20 µm).
transcript expression of ESC markers such as ABCG2, Myc, KLF4, Nanog, Oct4, Rex1, and Sox2 as well as NC markers such as Nestin, FOXD3, HNK1, MSX1, p75NTR, Sox9, Sox10 and N-cadherin (Fig. 3C). Collectively, these results suggested that 2D Matrigel in MESCM + 5% FBS could not maintain the TM phenotype although it promoted cell expansion.

Because Du et al. have reported that CD73, CD90, CD166 and Bmi1 are valuable stem cell markers, we have also performed their expression in TM cells at different culture stages. Although CD90 is a multipotent stem cell marker for the other model system, its transcript has not been found in TM cells in our system. Though the transcripts of CD73, CD166 or Bmi1 have been found in TM cells expanded in our system, CD73 is upregulated from passage 1 to passage 7, CD166 is downregulated from passage 2 to passage 7, and Bmi1 is upregulated at passage 1 (Supplemental Fig. 2A). In addition, the expression of those markers is not affected by 3D Matrigel (Supplemental Fig. 2B).

**TM phenotype and progenitor status is reversed by 3D Matrigel.** The gradual loss of the in vivo phenotype on 2D Matrigel has also been noted during expansion of human limbal niche cells and such a loss can be reversed by reseeding cells on 3D Matrigel. To test this, we passaged $2 \times 10^4$ per cm$^2$ of P2 TM cells on 2D Matrigel as a control and 3D Matrigel with or without Noggin in MESCM + 5% FBS. Upon reseeding back to 3D Matrigel for 48 h, these cells formed spheres (Fig. 4C). RT-PCR disclosed significant upregulation of the transcript level of TM cell markers such as AQP1, CHI3L1, MGP and AnkG except stromal marker vimenitin (Vim) (Fig. 4A) as well as embryonic stem cell (ESC) and NC markers such as Nanog, Oct4, Sox10 and PDGFR$\beta$ (Fig. 4B). Such upregulation of TM, ESC and NC markers except TM marker CHI3L1, ABCG2, Myc, Nestin, p75NTR and N-cadherin was attenuated by addition of Noggin (Fig. 4B and C). Immunostaining showed nuclear localization of Oct4, Sox2, Nanog and KLF4 in TM cells cultured on 3D Matrigel, but not in cells seeded on 2D Matrigel (Fig. 4C). Addition of Noggin abolished nuclear localization of Oct4, Sox2, Nanog and KLF4 in cells seeded on 3D Matrigel (Fig. 4C).

**BMP signaling is activated on 3D Matrigel Culture.** Because addition of Noggin abolished the reversal effect of 3D Matrigel in upregulating expression of TM markers as well as ESC and NC markers (Fig. 4), we would like to confirm the involvement of BMP signaling, which is also involved in the reversal of the gradual phenotypic loss of human limbal niche cells by 3D Matrigel. Compared to the control cultured on 2D Matrigel, TM cells exhibited notable upregulation of BMP2, BMP4, and BMP6 when reseeded on 3D Matrigel (Fig. 5A). The upregulation of BMP ligands was coupled with upregulation of BMP receptor 2 (BMPR2) but downregulation of BMPR1B (Fig. 5A). Immunostaining showed nuclear localization of pSmad1/5/8 in TM cells seeded on 3D Matrigel but not on 2D Matrigel (Fig. 5B), supporting the activation of canonical BMP signaling in the former but
not the latter. Blocking of BMP signaling by Noggin abolished up-regulation of BMP2, BMP4, BMP6 and BMPR2 (Fig. 5A), blocked nuclear translocation of pSmad1/5/8 (Fig. 5B), and downregulated transcript expression of TM markers (Fig. 4A) and markers of ESCs and NCs (Fig. 4B) and immunostaining of ESC markers (Fig. 4C). These results confirmed that activation of BMP signaling was critical for maintaining the TM phenotype and progenitor status.

Expanded TM cells are multipotent progenitors. Because expanded TM cells also express markers of ESC and NC (Figs 1 and 4), we wondered whether they were multipotent. Using P3 cells cultured on 2D Matrigel as a control, we subjected them together with P2 cells reseeded on 3D Matrigel to a number of differentiation assays achieved by respective differentiation media. For comparison, normal human corneal endothelial cells (HCEC) were cultured and formed a monolayer of hexagonal cells expressing a typical immunostaining pattern for several markers such as acetyl-α-tubulin (ciliary pattern), p120-catenin (p120), β-catenin, N-cadherin, Zona occludens protein 1 (ZO-1) and Na⁺/K⁺-ATPase (all intercellular junctions) (Fig. 6A, top panel)22,23. Before induction, TM cells on both 2D Matrigel and 3D Matrigel were spindle shaped (Fig. 6A, Phase) and did not exhibit any of the aforementioned HCEC staining pattern (Fig. 6A). However, after induction, TM cells on both 2D Matrigel and 3D Matrigel turned into more hexagonal and expressed an immunostaining pattern similar to HCEC with positive expression of p120, β-catenin, N-cadherin, ZO-1, and Na⁺/K⁺-ATPase preferentially in intercellular junctions (Fig. 6A). The staining for acetyl-α-tubulin was also peri-membranous as contrasted to the central punctate staining at the center of the hexagon as reported in HCEC22–25. All cells did not show any positive staining to LEF1 and S100A4, which are markers of endothelium-mesenchymal transition22 (not shown), indicating that these cells were not transitioned to a pathological state. These results suggested that TM cells expanded on 2D Matrigel or 3D Matrigel still retained the plasticity to be differentiated into HCEC-like cells. In addition, TM cells expanded on both 2D and 3D Matrigel could be differentiated into adipocytes (Fig. 6B). Aggregates of TM cells cultured on 3D Matrigel could, but those cultured on 2D Matrigel could not be induced to chondrocytes as evidenced by Alcian Blue staining (Fig. 6B), which was confirmed by measuring OD at 450 nm (not shown)18. In contrast, TM cells on both 2D and 3D Matrigel could not be induced into keratocytes as shown by negative staining of keratocan (Fig. 6C), which is corneal stromal specific extracellular matrix8 and could not be induced into osteocytes (not shown).

Discussion

Human trabecular meshwork cells are endothelial-like cells in a connective tissue located close to the front of the eye. However, most of authors described the cell shape as “spindle” when the cells were cultured in vitro26–28. We have found that the morphology of TM primary culture was indeed spindle when scattered (Fig. 1). We have also noted that TM cells would be spindle after confluence when cultured on different substrates except 2D Matrigel plus ME SCM-based media (Fig. 1), suggesting that TM cells can keep their in vivo phenotype when cultured on 2D Matrigel in ME SCM-based media.

TM cells characteristically express Vim (a stromal cell marker), aquaporin-1 (AQP-1), matrix gla protein (MGP) and chitinase 3-like 1 (CHI3L1) and AnK29–34. Interestingly, there is also a population of progenitors in Schwalbe’s Ring, the translation area between the periphery of corneal endothelium and the anterior non-filtering portion of TM, evidenced to have the capability of forming free-floating neurospheres, a function associated with multipotent progenitors35 expressing Oct-3/4, nestin, and Sox23,5. It is unclear how the entire population of TM
Our results support the conclusion that there is a population of TM progenitors in the trabecular meshwork6, and others have always included serum in their culturing of TM cells 10–15. Although the exact mechanism remains unknown to explain the necessity of FBS to expand TM cells, we speculate that this might have something to do with the matrix rigidity, which influence the cell shape. Herein, our comparative analysis disclosed that 2D Matrigel, i.e., a solubilized basement membrane matrix containing laminin, collagen IV, heparin sulfate proteoglycans, entactin/nidogen, and a number of growth factors (for review, see ref.35), was better than collagen IV and fibronectin in order to maintain TM as cuboidal cells and express the normal phenotype of TM cells based on the TM markers such as AQP1, MGP, CHI3L1, and AnkG (Fig. 1). This result resembles our success in using 2D Matrigel to isolate and expand human limbal niche progenitors16–18,20,36 and suggests that the in vitro extracellular matrix components such as laminins, collagens, elastin, fibronectin, fibrillins, proteoglycans, and matricellular proteins (reviewed in ref.25) collectively play an important role in the maintenance of TM phenotype. Unlike human limbal niche progenitors where in vivo expansion can be achieved in serum-free MESCM16–18,20, successful expansion of TM cells on 2D Matrigel requires addition of 5% FBS (Figs 2 and S1). We chose 2D culture for expansion and 3D culture for restoration of the normal phenotype based on our prior work on human limbal niche cells16–18,20. As noted in our prior work, direct seeding cells on 3D Matrigel maintains the normal phenotype but cells cease proliferation, thus resulting in the failure of ex vivo expansion. Similar to human limbal niche cells, TM cells cannot be expanded when seeded directly on 3D Matrigel. The underlying reason for 2D Matrigel to help expand TM cells or limbal niche cells remains unknown. We speculate that this might have something to do with the matrix rigidity, which influence the cell shape. Although the exact mechanism remains unknown to explain the necessity of FBS to expand TM cells, others have always included serum in their culturing of TM cells10–15.

Interestingly, such a population of TM cells also express ESC markers such as KLF4, Oct4, Sox2, Nanog, CD73, CD166 and Bmi1 but not CD90 and NC markers such as p75NTR, Sox9, FoxD3 (Figs 1, 3 and 4, Supplemental Fig. 2). The discrepancy for expression of CD90 from Du et al.8 is probably due to difference of the culture system, for example, different culture substrates and media. In addition, the cells form spheres when reseeded in 3D Matrigel after digestion in MESCM (Fig. 4). These phenomena resemble the characteristics of induced pluripotent stem cells (iPSCs) expressing Oct4, Sox2, KLF4 and Myc (OSKM)18, and limbal niche progenitors expressing embryonic stem cells and neural crest markers such as Oct4, Sox2, SSEA4, and Nanog, Nestin, N-Cadherin, and CD3416,18,19, suggesting this population of the cells are a group of the cells with adult progenitor characteristics. Our results support the conclusion that there is a population of TM progenitors in the trabecular meshwork6, and such a population of the progenitors not only express ESC and NC but also TM markers.

However on 2D Matrigel, like human limbal niche cells16–18, expanded TM cells also gradually lost expression of TM markers and ESC and NC markers (Fig. 3) and such a loss was transient because it could be regained by seeding P3 cells on 3D Matrigel in MESCM + 5% FBS to yield sphere growth (Fig. 4). Interestingly, the expression of CD73, CD166 and Bmi1 is not affected by 3D Matrigel (Supplemental Fig. 2). We also attribute this difference from Du et al.8 to the different culture systems. These results collectively support the use of this in vitro culturing model system to expand TM cells. In 3D Matrigel, also like human limbal niche progenitors40, the recovery of TM phenotype as progenitor status depends on activation of canonical BMP signaling (Fig. 5) as evidenced by the nullifying effect by BMP inhibitor Noggin (Figs 4 and 5). Our study showed that 3D Matrigel activates canonical BMP signaling (not only promotes expression of BMP2, BMP4 and BMP6, but also inhibits expression of BMPR1B and BMPR2), and promotes nuclear accumulation of pSMAD1/5/8 (Fig. 5). This finding of activation
of BMP canonical signaling is also reported in human limbal niche cells, of which the culturing system does not include FBS. This finding strongly suggests that activation of BMP signaling is not due to the inclusion of FBS, which contains multiple growth factors and cytokines.

Expanded human limbal niche progenitor cells in 3D Matrigel have the potential of differentiation into angiogenesis/pericyte progenitors and exhibit a better potential than human bone marrow MSCs to differentiate into adipocytes, chondrocytes, and osteocytes. Although P3 human TM cells expanded on 3D Matrigel also showed overexpression of ESC and NC markers (Fig. 4) [similar to human limbal niche progenitors although not tested for angiogenesis/pericyte differentiation], they exhibited the potential to differentiate into adipocytes and chondrocytes but not osteocytes (Fig. 6). P3 TM cells cultured on 2D or 3D Matrigel could be differentiated into adipocytes, however only P3 TM cells cultured on 3D Matrigel could be differentiated into chondrocytes (Fig. 6). Du et al. reported that their expanded human TM cells can be differentiated into keratocyte-like cells. Herein, we noted that our expanded TM cells were not able to differentiate into keratocyte-like cells due to negative staining of keratocan but could be differentiated into human corneal endothelium-like cells as assessed by the morphology and expression of HCEC markers (Fig. 6) that have been reported by us. The discrepancy between our finding and that by Du et al’s work in the lack of keratocyte differentiation might also be owing to the use of a different expansion medium. Collectively, these results strongly suggest that we have developed a novel in vitro model system to expand TM progenitors that have the potential of being differentiated into different NC lineages. That is, we have optimized the method of expanding TM progenitor cells using the unique combination of collagenase A (reported previously by Stamer et al.) and 2D Matrigel + MESCM + 5% FBS for effective isolation and expansion of TM cells and 3D Matrigel for phenotype reversion of the expanded TM cells to that in vivo. Using this method, we are able to expand the small cuboidal TM progenitors expressing markers of TM, ESC, and NC cells effectively. Such expanded TM progenitors may be used to study pathogenesis of glaucoma and corneal endothelial dysfunction and may also have the potential to be deployed as a regenerative therapy for treating these blinding diseases.

Methods

Materials. Materials used for isolation, culture and relevant experiments of TM cells were listed in Supplementary Table S4. Assay IDs for Real-time PCR primers were presented in Supplementary Table S5. Sources of antibodies used for immunofluorescence Staining were shown in Supplementary Table S6.

Isolation and Expansion of Human Corneal Endothelial Cells and TM Cells. This research protocol was approved by Institutional Review Board of Tissue Tech Inc., Miami, Florida. Thirty-two corneoscleral rims from 18 to 60 years old donors were obtained from the Florida Lions Eye Bank (Miami, FL) and managed in accordance with the Declaration of Helsinki. The identities of these anonymous cadaver donors could not be identified. In brief, human corneal endothelial cells (HCEC) were isolated as reported. Under a dissecting microscope, the corneoscleral rim was stripped off the adherent iris and was cut through the inner edge of Schwalbe’s line. After rinsing three times with MESCM, the Descemet’s membrane was stripped and digested at 37 °C for 16 h with 2 mg/ml collagenase A in MESCM + 5%FBS. The resultant HCEC aggregates were collected by centrifugation at 1,800 rpm for 5 min to remove the digestion solution and cultured in 24-well dishes coated with collagen IV in MESCM, which was made of DMEM/F-12 (1:1) supplemented with 10% knockout serum, 5 µg/ml insulin, 5 µg/ml transferrin, 5 ng/ml sodium selenite, 4 ng/ml Fibroblast growth factor-basic (bFGF), 10 ng/ml human leukemia inhibitory factor (hILF), 50 µg/ml gentamicin, and 1.25 µg/ml amphotericin. TM cells were isolated according to Du et al. and Stamer et al. with modifications. The remaining corneoscleral rim, after a cut made at the Schwalbe’s line, TM tissue was removed and digested with collagenase A in MESCM + 5% FBS at 37 ºC for 16 h. The resultant cells were washed once in MESCM + 5% FBS and seeded at a density of 2 × 10^4 per cm^2 on plastic or plastic coated with 2D Matrigel by adding 5% diluted Matrigel after 1 h incubation at 37 ºC, or coated with fibronectin (0.01% fibronectin) or collagen IV (10 µg/ml) after overnight incubation at 4 ºC in MESCM + 5% FBS, MESCM, or SCGM which is made of OptiMEM-1 supplemented with 5% fetal bovine serum, 10 ng/ml epidermal growth factor (EGF), 100 µg/ml bovine pituitary extract, 20 µg/ml ascorbic acid, 200 µg/ml calcium chloride, 0.08% chondroitin sulfate, 100 IU/ml penicillin, 100 µg/ml streptomycin, and 50 µg/ml gentamicin. Upon 80–90% confluence, cells on 2D Matrigel in MESCM or MESCM + 5%FBS were passaged serially at a density of 5 × 10^3 per cm^2. The extent of total expansion was measured by the number of population doubling using the following formula: number of cell doublings (NCD) = log_{10}(y/x)/log_{10}2, where y is the final density of the cells and x is the initial seeding density of the cells.

Maintenance of TM Progenitor Cells. P2 TM cells cultured on 2D Matrigel in MESCM + 5% FBS were passaged at the density of 5 × 10^4 cells per cm^2 on 2D Matrigel in MESCM + 5% FBS for 1 week or at the density of 1 × 10^5 cells per cm^2 on 3D Matrigel that was made by adding 50% diluted Matrigel in MESCM + 5% FBS for 48 h. The resultant P3 cells on either 2D or 3D Matrigel were then harvested for the following experiments. Spheres formed by TM cells on 3D Matrigel were isolated by digestion with 10 mg/ml dispase II at 37 °C for 2 h and rendered into single cells by 0.05% trypsin and 1 mM EDTA before being examined for their multipotent potential via various differentiation assays (see below) when compared to P3 cells cultured on 2D Matrigel. In addition, Noggin, a BMP signaling inhibitor, was also added at the final concentration of 500 ng/ml for 48 h in sphere cultures on 3D Matrigel.

Differential Assays. For differentiation into keratocytes, 3 × 10^4 cells were spun down for pellet preparation and cultured in the Keratocyte Differentiation Medium consisting of Advanced DMEM with 10 ng/ml fibroblast growth factor 2 and 0.5 mM ascorbic acid; Thermo Fisher Scientific) for 21 days. For differentiation into corneal endothelial cells, single cells were seeded at the density of 1 × 10^4 cells per cm^2 in 24-well plate coated with...
collagen IV in MESCM + 5% FBS. After cell adhesion in 24 h, the medium was switched to Corneal Endothelial Differentiation Medium consisting of low-glucose DMEM with 10% FBS and 50 μg/ml gentamicin, and 1.25 μg/ml amphotericin B for 28 days. For differentiation into adipocytes or osteocytes, single cells were seeded at the density of 1 × 10^4 cells per cm² in 24-well plate coated with Matrigel in MESCM + 5% FBS. After cell adhesion in 24 h, the medium was switched to the Adipogenesis Differentiation Medium (Invitrogen) or Osteogenesis Differentiation Medium (Invitrogen), respectively, for 21 days. For differentiation into chondrocytes, 3 × 10^5 cells were prepared for pellets and cultured in Chondrogenesis Differentiation Medium (Life Line) for 28 days. All the above media were changed every 3 days.

RNA Extraction, Reverse Transcription, and Real-time PCR. Total RNAs were extracted using an RNeasy Mini kit (QIAGEN) and reverse-transcribed using a High Capacity Reverse Transcription kit (Life Technologies). cDNA of each cell component was amplified by RT-real-time PCR using specific primer–probe mixtures and DNA polymerase in a real-time PCR system (7000; Life Technologies). The real-time RT-PCR profile consisted of 10 min of initial activation at 95°C, followed by 40 cycles of 15 sec denaturation at 95°C, and 1 min annealing and extension at 60°C. All TagMan Gene Expression Assays are listed in Supplementary Table S2.

Immunofluorescence, Oil Red O and Alcian Blue staining. For characterization of induction of endothelium, immunostaining of corneal endothelial markers such as acetyl-α-tubulin, p120, Na+/K+-ATPase, ZO-1, N-cadherin and β-catenin and endothelial-mesenchymal transition (EMT) markers such as LEF1 and S100A4 was performed as described. For characterization of induction of keratocytes, immunostaining of keratocyte markers keratan sulfate and keratocan was performed as reported. In brief, single cells were prepared by cytospin after tryptic/EDTA digestion and centrifuge at 1000 rpm for 8 min (StatSpin, Inc., Norwood, MA), fixed with 4% paraformaldehyde for 15 min, permeabilized with 0.2% Triton X-100 in PBS for 30 min, and blocked with 2% BSA in PBS for 1 h before being incubated with specific primary antibodies overnight at 4°C. After washing with PBS, samples were incubated with corresponding secondary antibodies for 1 h in room temperature using appropriate isotype-matched non-specific IgG antibodies as controls. The nucleus was counterstained with Hoechst 33342 before being analyzed with a Zeiss LSM 710 confocal microscope (LSM700; Carl Zeiss, Thornwood, NY). For characterization of induced adipocytes, the samples were stained with Oil Red O. Oil Red O-positive cells were counted in a total of 2000 cells in triplicates as reported. For characterization of induced chondrocytes, the samples were stained with Alcian Blue and quantified following the manufacturer’s protocol by measuring OD at 450 nm.

Statistical Analysis. All summary data were reported as means ± SD calculated for each group and compared using ANOVA and the Student’s paired or unpaired t test using Excel software (Microsoft). Test results were reported as two-tailed p-values, where p < 0.05 was considered statistically significant.

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Author Contributions
Y. Zhang and S.B. Cai have contributed to experimental design, performance of experiments. S.C.G. Tseng and Y.T. Zhu have overseen the experimental design, organized data and written the manuscripts.

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