The Silk-protein Sericin Induces Rapid Melanization of Cultured Primary Human Retinal Pigment Epithelial Cells by Activating the NF-κB Pathway

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Restoration of the retinal pigment epithelial (RPE) cells to prevent further loss of vision in patients with age-related macular degeneration represents a promising novel treatment modality. Development of RPE transplants, however, requires up to 3 months of cell differentiation. We explored whether the silk protein sericin can induce maturation of primary human retinal pigment epithelial (hRPE) cells. Microarray analysis demonstrated that sericin up-regulated RPE-associated transcripts (RPE65 and CRALBP). Upstream analysis identified the NF-κB pathway as one of the top sericin-induced regulators. ELISA confirmed that sericin stimulates the main NF-κB pathway. Increased levels of RPE-associated proteins (RPE65 and the pigment melanin) in the sericin-supplemented cultures were confirmed by western blot, spectrophotometry and transmission electron microscopy. Sericin also increased cell density and reduced cell death following serum starvation in culture. Inclusion of NF-κB agonists and antagonists in the culture medium showed that activation of the NF-κB pathway appears to be necessary, but not sufficient, for sericin-induced RPE pigmentation. We conclude that sericin promotes pigmentation of cultured primary hRPE cells by activating the main NF-κB pathway. Sericin’s potential role in culture protocols for rapid differentiation of hRPE cells derived from embryonic or induced pluripotent stem cells should be investigated.

Viable retinal pigment epithelial (RPE) cells are critical for normal photoreceptor cell metabolism and visual cycle. Hence, RPE degeneration is accompanied by concomitant photoreceptor degeneration1. Drusen, which are located between the RPE cells and Bruch’s membranes, contain a number of proteins that relate to inflammation; and there is increasing body of evidence that oxidative stress2, local inflammation3, and activation of the complement system4 play a part in the development of AMD. The choriocapillaris has also been suggested to play a role in the etiology of AMD5. The disorder is subdivided into a dry and a wet type. There are no established curative treatment options for the dry type of AMD, which constitutes more than 85% of the cases. Cell based restoration of RPE cells has been explored in several studies6,7. There are two main approaches for delivering cultured RPE to the submacular space: 1) injection of a RPE cell suspension; and 2) implantation of RPE as an intact cell sheet8. Benefits of the former include ease of procedure, while some of the disadvantages are increased apoptosis due to loss of cell-cell interaction and a disorganized RPE upon injection8. Implantation of RPE as an intact cell sheet should in theory prevent the mentioned disadvantages of injecting RPE.

Several culture protocols intended for driving the differentiation of human embryonic stem cells (hESC) and human induced pluripotent stem cells (iPSC) into RPE have been described9–12. Although protocols using hESC or iPSC have successfully produced differentiated and pigmented RPE cells, they are usually time-consuming,
sometimes necessitating a total culture period of up to 3 months\(^1^3\). Cutting down the time necessary for producing differentiated RPE reduces costs of production and lessens the risk of infection during prolonged cultures.

The silk protein sericin mainly consist of three polypeptides having molecular masses of 400, 250, and 150 kDa\(^1^4\) and encompasses 25–30% of all silk proteins\(^1^5\). It has been recognized to increase cell proliferation in serum-free media\(^1^6\), promote antioxidant effects\(^1^7\), inhibit the melanogenesis enzyme tyrosinase\(^1^7\), enhance wound healing\(^1^8\) and support the growth and attachment of fibroblasts\(^1^8\).

Sericin has demonstrated serum-sparing effects in media for cryopreservation of human dermal fibroblasts, human epidermal keratinocytes and rat islets\(^1^9,2^0\). The prospect of avoiding serum, either autologous or animal-derived, has become increasingly appealing with steadily tougher regulatory demands in regenerative medicine over the past years. Replacing serum with sericin could dramatically reduce the risk of infections, which for animal derived components may be particularly severe, not only for the patient, but also for the larger community\(^2^1\). Despite the rare occurrence of infections following the use of fetal bovine serum (FBS), measures taken to avoid both the risk of infections and standardize the culture protocols are most welcomed by regulatory authorities. In the current study, we explored the potential of using the silk protein sericin as serum-substitute in culture medium to induce maturation of primary human retinal pigment epithelial cells (hRPE). Our results revealed a surprising effect of sericin on promoting hRPE maturation by activating the NF-κB pathway.

**Results**

**Microarray Analysis of hRPE Cultured With or Without Sericin.** Microarray experiments were performed to assess the effect on hRPE of adding sericin to the culture medium in absence of FBS. Similarly as described in materials and methods, hRPE were seeded in complete EpiCM on Nunclon \(\Delta\) surface plates and cultured for two days before replacing EpiCM with either 1) DMEM with 1% sericin or 2) DMEM. Both DMEM media were supplied with 10.000 U penicillin and 10 mg streptomycin at a final concentration of 1%.

**Global Perspective.** Gene expression differed considerably between the two culture groups, with a total of 438 significantly differentially regulated genes (fold change > 1.5; \(P < 0.05\)). In the sericin-supplemented group, 229 genes were down-regulated and 209 genes were up-regulated compared to the control, while 14419 transcripts were unchanged.

**Upstream analysis.** The top upstream regulators were tumor necrosis factor (TNF) (z-score: 5.008), interferon-γ (IFN-γ) (z-score: 4.623), the NF-κB complex (z-score: 4.597) and interleukin-1β (IL1β) (z-score: 4.567). Pathway analysis of NF-κB revealed up-regulation of transcripts belonging to both the main and alternate NF-κB pathway (Fig. 1). Activation of the NF-κB pathway was indicated by up-regulation of several of its member transcripts, including TNFAIP3, MAP3K8 and NFKB2 (Table 1).

**Substantially Regulated Genes.** The C-X-C motif chemokine 10 (CXCL10, also known as IP-10; interferon gamma-induced protein 10) was the most up-regulated gene in the dataset, with a 55.5-fold up-regulation in the sericin-supplemented group compared to the control (Table 2). Complement component 3 (C3) was the second most up-regulated gene in the sericin group, with a 34.7-fold up-regulation. The chemokine (C-C motif) ligand 2 (CCL2) was up-regulated 21.6-fold in the sericin-supplemented group (Table 2). TNFAIP3 (tumor necrosis factor, alpha-induced protein 3; A20), which is involved in NF-κB regulation, was up-regulated 10.4-fold in the sericin group.

**Downstream Effects.** The results obtained from the Ingenuity Pathway Analysis (IPA) predicted a downstream effect compatible with increased cell viability and survival in the sericin-supplemented group (Fig. 2). In the cell viability category, 37 of 57 genes had an expression direction consistent with increased cell viability, yielding a Z-score of 2.8. In the cell survival category, 38 of 61 genes had an expression direction consistent with increased cell survival, yielding a Z-score of 2.7.

**RPE Related Gene Transcripts.** Pigmentation. The (IPA) detected 11 differentially expressed genes related to cell pigmentation (Table 3). Ten of the genes were up-regulated and one was down-regulated in hRPE cells cultured in DMEM with sericin compared to cells cultured in DMEM without sericin.

**Visual Cycle.** Several visual cycle genes were significantly up-regulated in hRPE cells cultured in DMEM with sericin compared to that of cells cultured in DMEM without sericin, while no visual cycle genes were significantly down-regulated (Table 4). Compared to cells cultured in DMEM without sericin, the cells that were cultured in DMEM with sericin displayed a 2.8-fold up-regulation of RPE65, a key isomerase of the visual cycle and an essential marker of RPE cell maturation\(^2^2,2^3\), and a 1.4-fold up-regulation of cellular retinaldehyde binding protein 1 (RBP1; also known as CRALBP), which positions retinol for enzymatic turnover and thereby accelerates the process\(^2^4\. Both retinol dehydrogenase 10 (RDH10) and retinol dehydrogenase 11 (RDH11) play complementary roles as 11-cis-retinol dehydrogenases in the visual cycle\(^2^5-2^7\), and were up-regulated 1.9 and 1.6-fold, respectively, in cells cultured in DMEM with sericin compared to cells cultured in DMEM without sericin.

**RT-PCR Validation.** To validate the microarray results, NFKBIA, RPE65, CSF1, NFKBIZ and RGR transcripts were quantified by RT-PCR in hRPE cells cultured in DMEM with or without 1% sericin for a total of 14 days. In Table 5, ΔΔCt values are transformed to fold change. All six transcripts were up-regulated in the Affymetrix and RT-PCR experiments (Table 5).
Microstructure Following hRPE Culture With or Without Sericin. Human retinal pigment epithelial cells were cultured with or without 1% sericin in two different basal media with or without FBS for 12 days before being assessed with light microscope for morphology and pigmentation. Widespread pigmentation was only seen in cells cultured with 1% sericin (DMEM/sericin; Fig. 3b, MEM-α/sericin; Fig. 3e), irrespective of the basal medium used (MEM-α or DMEM). Pigmented cells were rarely found in the groups that were cultivated without sericin (DMEM/FBS; Fig. 3a, DMEM; Fig. 3c, MEM-α/FBS; Fig. 3d, MEM-α; Fig. 3f). Typical cobblestone morphology with essentially hexagonal cells was seen in the samples cultured in DMEM with sericin (Fig. 3b), MEM-α with sericin (Fig. 3e), MEM-α with FBS (Fig. 3d), and MEM-α alone (Fig. 3f). Cell confluence was superior in DMEM with 1% sericin (Fig. 3b) compared to DMEM alone (Fig. 3c) and DMEM with 1% FBS (Fig. 3a). Hence, these results show that after 12 days in culture, only cells cultured in media supplied with sericin become pigmented and sericin appears to support hRPE cell confluence.

Ultrastructure Following hRPE Culture With or Without Sericin. Scanning electron microscopy of hRPE cultured for 12 days in DMEM with 1% sericin appeared tightly adjoined, hexagonal and had apical microvilli (Fig. 3g), hence demonstrating a differentiated ultrastructure. Transmission electron microscopy of hRPE cultured for 12 days in DMEM with 1% sericin contained melanosomes of all four stages following seven days of culture (Fig. 3h), thereby supporting the validity of the light microscopy experiments.

Melanin Quantification Following hRPE Culture With or Without Sericin. As the pigment melanin absorbs light at a specific wave length, measurement of pigment quantity is commonly performed by spectrophotometry\(^2^\). Following 12-day culture, spectrophotometry showed increased absorption at 562 nm in cells cultured...
in DMEM with 1% sericin (39 ± 6 melanin/protein; \( P < 0.001 \)) compared to cells cultured in DMEM without sericin (Fig. 4). Thus, the results further supports increased pigmentation in sericin-cultured cells.

Protein Levels of CRALBP and RPE65 Following hRPE Culture With or Without Sericin. To verify the presence and to assess the quantity of RPE-related proteins quantitative immunofluorescence and western blot experiments were used to analyze the mean levels of CRALBP and RPE65. With quantitative immunofluorescence, CRALBP was most abundant in cells cultured for 14 days in DMEM with sericin, and present to a significantly higher degree than in cells cultured in DMEM alone, DMEM with FBS or DMEM with FBS and sericin (\( P < 0.01 \)) (Fig. 5). Retinal pigment epithelial cells cultured in DMEM with sericin and FBS also demonstrated a significantly higher expression of CRALBP compared to the DMEM alone and DMEM with FBS groups (\( P < 0.01 \)). Thus, quantitative immunofluorescence confirmed increased levels of CRALBP, as indicated by the microarray results (Table 4). On western blot experiments, however, the level of CRALBP protein in cells cultured in DMEM with sericin (87% ± 21%) relative to that in cells cultured in DMEM alone (100%; \( P = 0.38 \)) was similar. The level of RPE65 protein, on the other hand, was significantly higher in the DMEM with sericin group (168% ± 11%) compared to in DMEM alone (100%; \( P = 0.008 \)), as measured by western blot. Therefore, sericin increases RPE65 protein in cultured primary hRPE.

hRPE Cell Density and Cell Death Following Serum Starvation. As the microarray results predicted increased viability of culturing cells in DMEM with sericin compared to DMEM without sericin, cell density and

| Gene Symbol | Gene Name | Affymetrix ID | FC | P-value |
|-------------|-----------|---------------|----|---------|
| TNFAIP3     | Tumor necrosis factor, alpha-induced protein 3 | 17012946 | 10.4 | 8.7E-05 |
| MAP3K8      | Mitogen-activated protein kinase kinase kinase 8 | 16703642 | 2.5 | 1.1E-02 |
| NFKB2       | Nuclear factor of kappa light polypeptide gene enhancer in B-cells 2 (p49/p100) | 16708623 | 2.1 | 3.3E-05 |
| TNFSF13B    | Tumor necrosis factor (ligand) superfamily, member 13b | 16776339 | 2.1 | 1.1E-03 |
| NFKB1       | Nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 | 16969300 | 2.0 | 2.3E-04 |
| TNFAIP3     | Tumor necrosis factor, alpha-induced protein 3 | 17012946 | 1.8 | 8.7E-05 |
| TNIP1       | TNFAIP3 interacting protein 1 | 17001763 | 1.6 | 3.5E-04 |
| RELB        | V-rel avian reticuloendotheliosis viral oncogene homolog B | 16863168 | 1.6 | 9.0E-04 |
| EGF         | Epidermal growth factor | 16969729 | 1.4 | 1.1E-02 |
| MAP3K1      | Mitogen-activated protein kinase kinase kinase 1, E3 ubiquitin protein ligase | 16984945 | 1.2 | 1.5E-02 |
| TGF alpha   | Transforming growth factor, alpha | 16898788 | 1.1 | 4.5E-02 |

Table 1. Differentially Expressed Genes in the NF\( \kappa \)-B-pathway in hRPE Cultured in DMEM with Sericin. hRPE = human retinal pigment epithelial cells; DMEM = Dulbecco’s modified eagle medium; FC = fold change.

| Gene Symbol | Gene Name | Affymetrix ID | FC | P-value |
|-------------|-----------|---------------|----|---------|
| CXCL10      | Chemokine (C-X-C motif) ligand 10 | 16977052 | 55.5 | 4.2E-06 |
| C3          | Complement component 3 | 16687784 | 34.7 | 1.2E-06 |
| CCL2        | Chemokine (C-C motif) ligand 2 | 16833204 | 21.6 | 9.3E-07 |
| IL6         | Interleukin 6 | 17044177 | 15.6 | 8.8E-06 |
| TNFAIP3     | Tumor necrosis factor, alpha-induced protein 3 | 17012946 | 10.4 | 8.7E-05 |
| ICAM1       | Intercellular adhesion molecule 1 | 16858137 | 9.1 | 2.0E-06 |
| PTX3        | Pentraxin 3, long | 16947357 | 8.0 | 1.4E-04 |
| ADA         | Adenosine deaminase | 16919466 | 7.7 | 2.6E-06 |
| EDNRB       | Endothelin receptor type B | 16779958 | 7.5 | 1.6E-05 |
| CNTNAP1     | Contactin associated protein 1 | 16834409 | 7.3 | 5.3E-06 |
| CLGN        | Calmegin | 16980051 | −5.3 | 4.0E-06 |
| ADAM28      | ADAM metallopeptidase domain 28 | 17066921 | −5.5 | 2.8E-05 |
| LUFZ2       | Leucine zipper protein 2 | 16722987 | −5.6 | 8.7E-05 |
| VCAN        | Versican | 16986913 | −6.4 | 2.6E-05 |
| MFP4        | Microfibrillar-associated protein 4 | 16842266 | −6.6 | 3.9E-06 |
| LRRC15      | Leucine rich repeat containing 15 | 16962911 | −6.9 | 3.2E-04 |
| ST8SIA4     | ST8 alpha-N-acetyl-neuraminide alpha-2,8-sialyltransferase 4 | 16998532 | −6.9 | 7.3E-06 |
| COL1A1      | Collagen, type XIV, alpha 1 | 17072162 | −9.2 | 4.5E-07 |
| SMOC2       | SPARC related modular calcium binding 2 | 17041798 | −9.8 | 2.8E-08 |
| GRIK3       | Glutamate receptor, ionotropic, kainate 3 | 16685330 | −16.4 | 5.3E-06 |

Table 2. Top Ten Up- and Down-regulated Genes in hRPE Cultured in DMEM with Sericin. hRPE = human retinal pigment epithelial cells; DMEM = Dulbecco’s modified eagle medium; FC = fold change.
cell death after 12 days of cultivation with or without sericin was quantified by counting DAPI-stained cell nuclei and by microplate fluorometer measurements of EH-1, the latter which is indicative of dead cells. DMEM with sericin yielded the highest cell density and significantly higher density than DMEM alone ($P = 0.006$), DMEM with FBS ($P < 0.001$) and DMEM with sericin and FBS ($P = 0.007$) (Fig. 6a).

Cultured hRPE were assayed with EH-1 to quantify the number of dead cells following 14 days of culture. The highest number of dead cells was obtained when using DMEM alone (Fig. 6b). Cells cultured in this medium

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### Table 3. Differentially Expressed Pigmentation-associated Genes in hRPE Cultured in DMEM with Sericin.

| Gene Symbol | Gene Name                                      | Affymetrix ID   | FC   | P-value   |
|-------------|------------------------------------------------|-----------------|------|-----------|
| IL6         | Interleukin 6                                  | 17044177        | 15.6 | 8.8E-06   |
| EDNREB      | Endothelin receptor type B                     | 16779958        | 7.5  | 1.6E-05   |
| SOD2        | Superoxide dismutase 2, mitochondrial           | 17025267        | 4.6  | 1.6E-05   |
| C1orf11     | Chromosome 10 open reading frame 11            | 16706350        | 3.3  | 1.1E-05   |
| IRF1        | Interferon regulatory factor 1                 | 16999776        | 3.0  | 1.0E-03   |
| PROM1       | Prominin 1                                     | 16974534        | 2.9  | 2.6E-05   |
| INSIG1      | Insulin induced gene 1                         | 17053892        | 2.9  | 1.2E-04   |
| SLC24A5     | Solute carrier family 24, member 5             | 16800764        | 2.2  | 1.8E-04   |
| SOD3        | Superoxide dismutase 3, extracellular          | 16965519        | 2.1  | 4.1E-04   |
| TGM2GF2     | Transglutaminase 2                             | 16919158        | 2.0  | 6.9E-04   |
| HLLS        | Helicase, lymphoid-specific                     | 16707695        | 2.0  | 3.4E-03   |

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**Figure 2. Downstream effect upon culturing of human retinal pigment epithelial cells with or without sericin.** Ingenuity Pathway Analysis predicted a downstream effect compatible with increased viability and survival of human retinal pigment epithelial cells cultured in DMEM with 1% sericin compared to cells cultured without sericin. In the cell viability category (right center), 37 of 57 genes had an expression direction consistent with increased cell viability, yielding a Z-score of 2.8. In the cell survival category (left center), 38 of 61 genes had an expression direction consistent with increased cell survival, yielding a Z-score of 2.7. Red symbols indicate increased transcript levels upon culturing in presence of sericin, while green symbol indicate reduced transcript levels. The dotted lines indicate indirect relationships leading to activation (orange) or inhibition (blue). Yellow and grey dotted lines indicate inconsistent relationships and no predicted effects, respectively.
Necessary, but not sufficient, for sericin-induced melanization of hRPE. B activation appears to be of DMEM without sericin did not induce melanization (Figs. 4 and 7c). Thus, NF-κB activation, as opposed to control cells cultured in DMEM with sericin, but without JSH-23 (Figs. 4 and 7a) (N.

Development of pigmented cells as demonstrated by light microscopy and spectrophotometry following seven days of cultures. Data are average from triplicates with MAPRE1 and POLR5H as endogenous controls. RT PCR analysis demonstrated a significantly higher amount of dead cells than when using DMEM with sericin or DMEM with FBS (P < 0.001). Thus, the cell density and cell death results supported the microarray prediction of increased viability when culturing hRPE in DMEM with sericin.

**NF-κB Pathway and Melanization of Cultured hRPE Cells.** Since IPA identified both the NF-κB complex, and TNF, a well known activator of NF-κB, as top upstream regulators, we focused on the role of NF-κB signaling in melanization. The NF-κB-inhibitor JSH-23 specifically inhibits nuclear translocation and activation of NF-κB. The addition of JSH-23 to a culture medium consisting of DMEM with 1% sericin prevented development of pigmented cells as demonstrated by light microscopy and spectrophotometry following seven days of culture, as opposed to control cells cultured in DMEM with sericin, but without JSH-23 (Figs. 4 and 7a,b) (N = 3). However, stimulation of the NF-κB pathway by adding PMA, a NF-κB agonist, to a culture medium consisting of DMEM without sericin did not induce melanization (Figs. 4 and 7c). Thus, NF-κB activation appears to be necessary, but not sufficient, for sericin-induced melanization of hRPE.

**Effect of Sericin on the Main and Alternate NF-κB Pathways.** To verify the microarray data, which indicated that sericin induces activation of the NF-κB pathway, we next investigated sericin’s effect on activation of the main (phospho-NF-κB p65) and alternate (NF-κB p52) pathways. Following 12 days, primary hRPE cells cultured in DMEM with 1% sericin exhibited a higher level of phospho-NF-κB p65 divided by total protein (7.6 ± 0.7) compared to cells cultured in DMEM alone (5.9 ± 0.1; P = 0.01). To investigate if FBS inhibits sericin’s stimulatory effect on phospho-NF-κB p65, we next compared cells cultured in DMEM with 1% sericin with that of cells cultured in DMEM with 1% sericin and FBS. The addition of FBS caused an even higher level of phospho-NF-κB p65 (12.4 ± 1.7; P = 0.01) compared to the DMEM with 1% sericin group. Regarding the alternate pathway, the mean level of NF-κB p52 protein in cells grown in DMEM with 1% sericin (2.0 ± 0.6; P = 0.12) relative to that in cells grown in DMEM alone was approximately two-fold higher, but the potential difference between the groups did not reach significance (N = 3) (Fig. 8). Cells cultured in DMEM with 1% sericin and FBS (2.7 ± 0.6; P = 0.04) demonstrated a more than two-fold increase in the level of NF-κB p52 protein relative to cells grown in DMEM alone (Fig. 8). Collectively, these data verify that sericin activates the main NF-κB pathway and that FBS do not inhibit sericin’s activation.

**Discussion**

In the current study, sericin induced pigmentation of cultured hRPE by NF-κB pathway activation while still preserving cell viability and improving RPE maturation, as indicated by pathway analysis of Affymetrix microarray, micro and ultra-structural studies, spectrophotometry, western blot, ELISA, and viability assays. The NF-κB pathway was, alongside TNF, IFN-γ, and IL1β, one of the top upstream sericin-induced regulators in this study. The NF-κB pathway is involved in multiple cellular processes, including inflammation.

**Table 4. Differentially Expressed Visual Cycle-associated Genes in hRPE Cultured in DMEM with Sericin.** hRPE = human retinal pigment epithelial cells; DMEM = Dulbecco’s modified eagle medium; FC = fold change.

| Gene Symbol | Gene Name | Assay FC | Assay P-value | Affymetrix FC | Affymetrix P-value |
|-------------|-----------|----------|---------------|---------------|-------------------|
| NFKBIA      | Nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, alpha | 12.9 | 4.0E-06 | 6.9 | 2.0E-03 |
| RPE65       | Retinal pigment epithelium-specific protein 65kDa | 4.3 | 1.8E-07 | 2.8 | 2.7E-05 |
| CSF1        | Colony stimulating factor 1 (macrophage) | 6.8 | 1.8E-04 | 2.8 | 1.0E-02 |
| NFKBIZ      | Nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, zeta | 5.1 | 2.0E-03 | 2.6 | 2.0E-02 |
| RGR         | Retinal G protein coupled receptor | 3.6 | 5.4E-05 | 2.3 | 2.0E-03 |

**Table 5. Verification of Affymetrix data by RT PCR.** The table shows average fold change (increase) in mRNA levels upon incubation of human retinal pigment epithelial cells with sericin for 12 days as compared to controls. Data are average from triplicates with MAPRE1 and POLR5H as endogenous controls. RT PCR = real time polymerase chain reaction; FC = fold change.
Figure 3. Micro- and ultrastructure of human retinal pigment epithelial cells cultured with or without sericin. Photomicrographs (a–f) show human retinal pigment epithelial cells cultured for 12 days with or without 1% sericin or 10% fetal bovine serum (FBS) using two different basal media. (a) DMEM/FBS; (b) DMEM/sericin; (c) DMEM; (d) MEM-α/FBS; (e) MEM-α/sericin; (f) MEM-α. Cellular pigmentation is indicated by white arrowheads and cobblestone morphology is indicated with black arrowheads. Magnification: 200×. Photomicrographs were captured by a black-and-white camera and are representative of eight samples. Electron microscopy analyses (g,h) were performed on human retinal pigment epithelial (hRPE) cells cultured in DMEM with 1% sericin for 12 days. (g) Scanning electron microscopy image showing the apical surface of a confluent layer of hexagonal hRPE cells with extensive microvilli (arrow). Image is representative of three samples. (h) Transmission electron microscopy image showing a polarized hRPE cell with apical tight junctions (black arrows), a basal nuclei and numerous melanosomes between stage I (non-melanized) and stage IV (melanized). Image is representative of four samples.

Figure 4. Effect of sericin on melanin production. The bar chart shows levels of melanin adjusted for total protein after culturing human retinal pigment epithelial cells for seven days in DMEM with or without 1% sericin, the NF-κB activator inhibitor 4-methyl-N1-(3-phenylpropyl)-1,2-benzenediamine (JSH-23) and the NF-κB agonist phorbol-12-myristate-13-acetate (PMA). N = 3. Error bars: standard deviation. *P < 0.05 compared to all other groups.
and immunity. NF-κB is also part of the TNF pathway and is regulated by IL1ß. A link between inflammatory cytokines, including IL1ß and TNF-α, and induction of pigmentation in chick RPE cells has been reported. Interestingly, IFN-γ activation has been related to hypopigmentation in skin melanocytes. In our study, sericin activated the main NF-κB pathway. Furthermore, the addition of the NF-κB activator inhibitor JSH-23 prevented pigmentation in sericin-cultured cells and thereby confirmed the role of NF-κB in pigmentation of hRPE, whereas the relatively scarce pigmentation achieved with the NF-κB agonist PMA alone suggested that sericin promotes pigmentation, albeit not exclusively, by activating the NF-κB pathway.

Ingenuity pathway analysis identified several genes that are potentially linked to sericin-induced pigmentation, including PROM1, C10orf11, SLC24A5, TGM2 and IRF1, which were all up-regulated by sericin. PROM1 is also related to maculopathy, as mutations in this gene may cause macular degeneration, including dominant bull’s eye maculopathy. Mutations in C10orf11 have been shown to decrease pigmentation of melanocytes and lead to human albinism. Interestingly, down-regulation of SLC24A5 causes reduced melanin content in chick RPE. Furthermore, SLC24A5 has also been related to melanin content in skin melanocytes and the gene product of SLC24A5 localizes to intracellular membranes, including melanosomes. Inhibition of TGM2 has been reported to suppress melanogenesis in human melanoma cells. Surprisingly, up-regulation of IRF1 has been associated with hypo-pigmentation in skin melanocytes.

Previous reports have demonstrated that sericin inhibits tyrosinase, which is the main rate-limiting melanogenesis enzyme, thus the induction of pigmentation by sericin in our study is unexpected. Tyrosinase catalyses the formation of dihydroxyphenylalanine (DOPA) from L-tyrosine. DOPA is subsequently converted to melanin, aided by tyrosinase-related proteins 1 and 2 (TRP-1 and TRP-2). Tyrosinase is inhibited by acidic conditions, including those resulting from high metabolic activity. The microarray data did not reveal any significant effect of sericin on the tyrosinase transcript (TYR), or on TRP-1 and TRP-2 encoding genes, (TYRP1 and TYRP2, respectively), thereby suggesting that sericin’s effect on pigmentation is unrelated to regulation of these genes. The production of melanin is a complex process, however, involving several steps where sericin could have a stimulating role that fully compensates for its acclaimed tyrosinase-inhibiting effect.

Pigmentation was almost exclusively seen in cells that had been cultured in sericin-supplemented basal media (either MEM-α or DMEM). Hexagonal cobblestone morphology was also achieved when using either MEM-α or DMEM supplemented with sericin. After culturing the cell line ARPE-19 for 98 days in DMEM with FBS, Ahmado and co-workers demonstrated pigmented and hexagonal cells. To our knowledge, this medium has
not been used for normal RPE. The MEM-α-based medium, however, has been shown to induce pigmentation in normal hRPE cells in study by Sonoda et al.44. In contrast to our study, the hRPE in their study was pigmented upon start of culture and, after initial depigmentation, became repigmented after 14 days. Both MEM-α and DMEM supplemented with FBS are known to induce maturation of RPE cells in prolonged culture (>3 weeks44/months43). Our culture time with sericin of seven to 14 days may, therefore, have been too short time for widespread melanogenesis to occur in these media.

In RPE, pigment can be seen within vesicles called melanosomes, which can be divided into four stages of development based on ultrastructure45. While stage I and II melanosomes are amelanotic, stage III and IV melanosomes are partially, and fully, pigmented, respectively. hRPE cultured in DMEM containing sericin displayed melanosomes of all four stages following 14 days of culture, which suggests that the process of melanogenesis was ongoing.

The tight junction barrier is involved in creating a polarized epithelium and is necessary for maintaining an apical-basal concentration gradient across the RPE45. Thus, as shown by TEM in the current study, sericin enabled the development of a polarized RPE, as indicated by presence of apical tight junctions and basal cell nuclei. 

**Figure 6. Effect of sericin on cell survival.** To assess the effect of sericin on cell survival confluent layers of human retinal pigment epithelial cells were cultured in DMEM with or without sericin or fetal bovine serum (FBS) for 12 days. (a) Bar chart showing cell density in fold change relative to the DMEM-group. Cell density was measured by counting DAPI-stained nuclei using ImageJ. N = 4 to 8. Error bars: standard deviation. *P < 0.01 compared to all other groups. (b) Bar chart showing amount of dead cells in fold change relative to the DMEM-group. Cell death was measured by ethidium homodimer-1 (ETH-1) uptake by dead cells using a microplate reader. N = 8. Error bars: standard deviation. *P < 0.001 compared to the other two groups.
the anti-angiogenic effect of CXCL10. CCL2 contributes in maintaining normal RPE morphology, and lack of the gene leads to RPE cell loss and stress56.

Inclusion of sericin in the culture medium leads to up-regulation of several genes related to the visual cycle, including RPE65, RDH10 and CRALBP. Retinal diseases can result from mutations or malfunction of key proteins in the visual cycle, in which the RPE serves as a critical component24. The RDH10 is essential for synthesis of embryonic retinoic acid and therefore for limb, craniofacial and organ development57. RPE65 is one of the key markers of RPE cells and is responsible for light-independent conversion of all-trans-retinyl esters into 11-cis-retinol58. The data for CRALBP, a marker of hRPE maturation that is involved in retinol recycling43,59, were inconclusive with respect to whether this protein was significantly changed by addition of sericin. RPE65, however, was upregulated by sericin, as indicated by microarray profiling, RT-PCR and western blot. Thus, sericin affects at least some of the key markers of the visual cycle in cultured primary hRPE.

Figure 7. Effect of stimulation/inhibition of NF-κB signaling. The NF-κB activator inhibitor 4-methyl-N1-(3-phenylpropyl)-1,2-benzenediamine (JSH-23) and the NF-κB agonist phorbol-12-myristate-13-acetate (PMA) was used to assess whether activation of the NF-κB pathway is necessary and sufficient for sericin-induced pigmentation of human retinal pigment epithelial (hRPE) cells following seven days of culture. (a) Photomicrograph showing pigment-containing hRPE cells (white arrows) following culture in DMEM with sericin. Magnification: 200×. (b) Photomicrograph showing absence of pigment in hRPE cells following culture in DMEM with sericin and JSH-23. Magnification: 200×. (c) Photomicrograph showing scarce pigment in hRPE cells (white arrow) following culture in DMEM with PMA. Magnification: 200×. Photomicrographs have been converted to gray scale images and are representative of three samples.
Viability analyses were performed to investigate whether the sericin-induced up-regulation of several inflammatory cytokines was accompanied with increased cell death. To reduce the effect of cell proliferation on cell density, the measurements of cell density were performed after 12 days of post-confluent culture with or without sericin or FBS. Sericin appeared to preserve cell density under serum-free conditions, and resulted in higher cell density than when adding either FBS or a combination of FBS and sericin to DMEM. Corroborating experiments with ETH-1 demonstrated that sericin promotes cell survival, which is in line with the down-stream prediction made by IPA. Our results are further supported by a study demonstrating that sericin protects against cell death following acute serum-deprivation and studies showing that FBS can be replaced by sericin in cryopreservation media without compromising viability. The increased survival of hRPE upon sericin may be related to up-regulation of TNAIP3, which inhibits TNF-induced apoptosis, or up-regulation of anti-oxidant genes, including the pigmentation-related gene SOD2. Downstream analysis by IPA also predicted a relationship between up-regulation of TNAIP3 and SOD2 and increased cell viability and cell survival. In addition, a direct reactive oxygen species-scavenging effect of sericin has been reported elsewhere. In RPE, inflammatory cytokines, such as IFN-γ and TNF-α, have been shown to induce SOD2, which promotes cell survival in the presence of oxidative stress. Thus, even though sericin promoted augmented expression of the inflammatory NF-κB pathway, cell survival was increased, possibly by the concomitant up-regulation of anti-apoptotic and anti-oxidant genes.

In conclusion, sericin promotes pigmentation of cultured hRPE by activating the main NF-κB pathway. Sericin's potential role in culture protocols for rapid differentiation of RPE cells derived from embryonic or induced pluripotent stem cells should be investigated.

**Methods**

Normal primary hRPE and complete epithelial cell medium (EpiCM) were obtained from ScienCell Research Laboratories (San Diego, CA). Dulbecco's modified eagle's medium (high glucose, with pyruvate; hereafter named DMEM), Minimal Essential Medium (α-modification; MEM-α), heat-inactivated fetal bovine serum (FBS), N1 growth supplement, taurine, triiodo-thyronine, non-essential amino acids, glutamine-penicillin-streptomycin, hydrocortisone, propidium iodide (PI) and 4′,6-diamidino-2-phenylindole (DAPI) and 4-methyl-N1-(3-phenylpropyl)-1,2-benzenediamine (JSH-23) were obtained from Sigma Aldrich (St Louis, MO).家系 Δ surface plates, pipettes and other routine plastics came from VWR International (West Chester, PA). Mouse monoclonal anti-cellular retinaldehyde-binding protein (CRALBP; clone B2) was from Abcam (Cambridge, UK). Ethidium homodimer-1 (ETH-1) was from Invitrogen. Secondary Cy3-conjugated anti-mouse antibody was purchased from Abcam (Cambridge, UK). Soluene®-350 was purchased from PerkinElmer (Waltham, MA). Pierce BCA Protein Assay Kit was obtained from Bio-Rad (Hercules, CA).

**Cell Culture.** Third passage hRPE were seeded (7000 cells/cm²) in complete EpiCM on Nuncelon Δ surface plates and cultured under routine conditions of 95% air and 5% CO₂ at 37 °C. A confluent layer was obtained after two days, at which time EpiCM was replaced with either: 1) DMEM with 1% FBS; 2) DMEM with 1% sericin; 3) DMEM without FBS or sericin; 4) MEM-α with 1% FBS; 5) MEM-α with 1% sericin; or 6) MEM-α without FBS or sericin. All culture media based on DMEM were supplied with 10,000 U penicillin and 10 mg streptomycin at a final concentration of 1%. The MEM-α-based media were added taurine, triiodo-thyronine, non-essential amino acids, glutamine-penicillin-streptomycin, hydrocortisone, and N1 medium supplement, as described elsewhere. The culture medium was changed every two to three days, and the hRPE were maintained in culture for 12 days. Bar chart shows amount of protein relative to the DMEM group. N = 3. *P = 0.04 compared to DMEM alone.
a total of between nine to 14 days (two days in EpiCM and the remaining days in the DMEM or MEM-α based media).

**RNA Extraction and Microarray Hybridization.** hRPE cells were cultured for 14 days and washed with PBS and lysed with QIAzol Lysis Reagent (N = 3). The lysate was transferred to a microcentrifuge vial. Total RNA was then purified according to the manufacturer's protocol, and 100 ng of total RNA was processed with a GeneChip HT One-Cycle cDNA Synthesis Kit and a GeneChip HT IVT Labeling Kit (Affymetrix, Santa Clara, CA). Labeled and fragmented single stranded cDNAs were hybridized to the GeneChip Human Gene 1.0 ST Arrays (28,869 transcripts) (Affymetrix). Thereafter, the arrays were rinsed and stained using a FS-450 fluidics station (Affymetrix). Signal intensities were measured with a Hewlett Packard Gene Array Scanner 3000 7G (Hewlett Packard, Palo Alto, CA), and the scanned images were processed by the Affymetrix GeneChip Command Console (AGCC). The CEL files were imported into Partek Genomics Suite software (Partek, Inc. MO, USA). Robust microarray analysis (RMA) was applied for normalization. Gene transcripts with a maximal signal value less than 32 across all arrays were removed to filter for low and non-expressed genes, reducing the number of gene transcripts to 23190. Differentially expressed genes between groups were identified using one-way ANOVA analysis in Partek Genomics Suite Software. Clustering analysis was made using the same name module in a Partek Genomics Suite Software. Gene transcripts with maximal signal values of less than 5 were removed to filter for low and non-expressed genes, resulting in 14419 gene transcripts. For expression comparisons of different groups, profiles were compared using a 1-way ANOVA model. Data were presented as fold changes (FC) and P-values.

**Microarray Data Analysis.** Upstream analysis, pathway analysis and downstream predictions were performed using Ingenuity Pathways Analysis (IPA) (www.ingenuity.com).

**Verification of Affymetrix Data by RT-PCR.** The differential gene expression data were validated for selected transcripts using TaqMan® Gene Expression Assays and the Applied Biosystems® ViiA™ 7 Real-Time PCR system (Applied Biosystem, Life technologies, Carlsbad, CA) (Table 6). Affymetrix analysis identified MAPRE1 and POLR5H as ideal for endogenous controls due to their stable expression across samples and groups. Briefly, 200 ng totalRNA were reverse transcribed using qScript™ cDNA Super Mix (Quanta Biosciences, Gaithersburg, MD) following the manufacturer’s instructions. After completion of cDNA synthesis 1/10 of the first strand reaction were used for PCR amplification. 9 μl of cDNA (diluted in H2O), 1 μl of selected primer/probes TaqMan® Gene Expression Assays (Life Technologies) and 10 μl TaqMan® Universal Master Mix (Life Technologies) following the manufacturer’s instructions. Normalized ΔCt values were calculated by subtracting the average Ct of the two endogenous controls from the Ct of the reference gene. ΔΔCt values were then calculated by subtracting normalized ΔCt values of the group containing cells cultured in DMEM with 1% sericin from the control group, which included cells cultured in DMEM without sericin (N = 3). P-values were calculated using Student's t-test in Microsoft Excel using delta Ct values. Normalized target gene expression levels (FC) were calculated using the formula: 2^(-ΔΔCt).

**Light Microscopy.** Cell morphology and presence of pigment was assessed after 14 days of culture by light microscopy at 200× magnification (N = 8). Photomicrographs were captured using a Leica DM IL LED microscope and Canon EOS 5D mark II camera or with a Nikon Eclipse microscope with a DS-Qi1 black-and-white camera.

**Transmission Electron Microscopy.** hRPE cells cultured for 12 days in DMEM with 1% sericin were processed for transmission electronmicroscopy (TEM) analysis as previously described64. In brief, ultrathin sections (60-70 nm thick) were cut on a Leica Ultracut Ultramicrotome (Leica, Wetzlar, Germany) and examined using a CM120 transmission electron microscope (Philips, Amsterdam, the Netherlands) (N = 4).

**Scanning Electron Microscopy.** hRPE cells cultured on glass coverslips in DMEM with 1% sericin for 12 days were used for scanning electron microscopy (SEM) as described previously65. Glutaraldehyde-fixed samples (N = 3) were dehydrated in increasing ethanol concentrations and then dried according to the critical point method (Polaron E3100 Critical Point Drier, Polaron Equipment Ltd., Watford, UK) with CO2 as the transitional fluid. The specimens were attached to carbon stubs and coated with a 30-nm thick layer of platinum in a Polaron Command Console (AGCC). The CEL files were imported into Partek Genomics Suite software (Partek, Inc. MO, USA).

| Gene symbol | Assay ID          |
|-------------|-------------------|
| RPE65       | Hs01071462_m1     |
| RGR         | Hs00173619_m1     |
| NFKBIA      | Hs00355671_g1     |
| NFKBIZ      | Hs00230071_m1     |
| CSF1        | Hs00174164_m1     |
| POLR3H      | Hs00978014_m1     |
| MAPRE1      | Hs01121102_g1     |

Table 6. Taqman Assays.
Melanin Measurement by Spectrophotometry. Intracellular melanin was quantified in hRPE cultured in DMEM with and without 1% serine, JSH-23 and PMA. Seven-day cultures in 25 cm² flasks were rinsed with PBS, and then re-suspended in 200 µL of RIPA lysis buffer consisting of 25 mM Tris-HCL [pH 7.6], 150 mM NaCl, 1% NP-40, 1% sodium deoxycholate and 0.1% sodium dodecyl sulfate dissolved in H₂O. Part of the cell lysates were then incubated with Pierce BCA protein assay kit for 30 minutes at 37 °C, and cooled to room temperature before measuring the absorbance at 562 nm using a microplate reader (VERSAmax, Molecular Devices, Sunnyvale, CA). Soluene®-350 was thereafter added to the remaining cell lysate (9:1). The cell lysate was subsequently incubated for 60 minutes at 80 °C, before being centrifuged for 10 minutes at 8600 g. Solubilized melanin was measured at 490 nm on the microplate reader, and the concentration was adjusted by multiplication with protein levels ($N_\text{protein}$).

Quantitative Immunofluorescence. Following 14 days of culture the cells were fixed in 100% methanol for 15 minutes and then washed three times with fresh PBS ($N = 4–8$). Fixed cells were incubated for 45 minutes at room temperature in a blocking buffer consisting of 10% goat serum, 1% BSA, 0.1% Triton X-100, 0.05% Tween-20, 0.05% sodium azide in PBS. Cells were then incubated overnight at 4 °C with the following antibody diluted in blocking buffer: anti-CRALBP (1:100), which targets a functional protein in the visual cycle and a marker for differentiated hRPE. The Cy3-conjugated secondary antibody, diluted in 0.2% PBST with 1% BSA, were incubated for one hour at room temperature. Negative control consisted of replacing the primary antibody with PBS. The cultures were thereafter rinsed three times in PBS and incubated with 1µg/mL DAPI in PBS to stain cell nuclei before a final wash with PBS.

Photomicrographs of the cultures were captured at 200 × magnification using a Nikon Eclipse Ti fluorescence microscope with a DS-Qi1 black-and-white camera. The exposure length and gain was maintained at a constant level for all samples, and the fluorescence intensities of the Cy3 fluorochromes, which were conjugated to the secondary antibodies, were within the dynamic range of the camera.

Phenotype was quantified using ImageJ (National Institutes of Health, Bethesda, MD) as described elsewhere[66,67], with some modifications. In brief, mean fluorescence per cell was measured by enlarging regions of interest (ROI) created around the DAPI-stained nuclei to enclose the CRALBP-expressing cytosol. By using this method, we were able to normalize for differences in cell density in each photomicrograph.

Quantification of Cell Density. After 14 days in culture, the hRPE ($N = 4–8$) were fixed with methanol, as described above. The cultures were then rinsed three times in PBS and incubated with 1 µg/mL DAPI in PBS to stain cell nuclei before a final wash with PBS. Photomicrographs of the cultures were captured at 200 × magnification with identical exposure length and gain. ImageJ was used to convert 16-bit images to binary images, as reported elsewhere[66]. The “Analyze particles” function in ImageJ was then used to automatically count cell nuclei per image.

Quantification of Cell Death. The amount of dead cells in the cultures after 14 days was quantified by incubating the samples with EH-1 for 30 minutes at 37 °C ($N = 8$). Ethidium homodimer-1 stains nuclei of dead cells red and its fluorescence was quantified by the microplate fluorometer with the excitation/emission filter pair 530/620. Background fluorescence, measured in wells incubated with EH-1-reagent, but without cells, was subtracted from all values before calculating mean fluorescence for the groups.

Western Blot. Protein levels were measured by Western blot analysis of primary hRPE cultured for 12 days. In brief, cells were lysed in RIPA-buffer and proteins analyzed by SDS-PAGE followed by electroblotting onto PVDF membranes.

The membranes were blocked with 5% bovine serum albumin/TBS at ambient temperature and incubated over-night at 4 °C with rabbit polyclonal anti-CRALBP (1:1500), mouse monoclonal anti-RPE65 (1:5000) or mouse monoclonal anti-NF-κB p100/p52 (1:4000). Samples were then incubated for one hour at ambient temperature with HRP-conjugated goat anti-rabbit (1:10.000) or goat anti-mouse (1:10.000) secondary antibodies. Immunoreactive bands were detected using enhanced chemiluminescence (ECL) substrate (Pierce Chemical Co, Rockford, USA), and visualized using the ImageQuant LAS4000 system (GE Healthcare, USA). To confirm the equality of protein lysates loading, membranes were subjected to mild stripping as described (http://www.abcam.com/ps/pdf/protocols/stripping%20for%20reprobing.pdf), and re-probed using a mouse anti-actin monoclonal antibody (1:3000), then processed as above. Intensity of the protein bands were quantified by ImageJ.

ELISA. Level of phospho-NF-κB p65 was measured by enzyme-linked immunosorbent assay (ELISA) in primary hRPE cultured for 12 days according to the manufacturers instructions (RayBio® Phospho-NF-κB p65 (S536) and Total NF-κB p65 ELISA kit). Briefly, samples were incubated with anti-phospho-NF-κB p65 for 2.5 hours at ambient temperature, washed, incubated with primary antibody for 1 hour, washed and then incubated with HRP-conjugated anti-rabbit IgG. Following washing, the addition of TMB substrate solution and Stop Solution, the color changes were measured at 450 nm.

Statistical Analysis. One-way ANOVA with Tukey’s (equal variances assumed) or Dunnett’s T3 (equal variances not assumed) post hoc pair-wise comparisons were used to compare three or more groups, while Student’s t-test was used to compare two groups (SPSS ver. 21.0). Data were expressed as mean ± standard deviation, and values were considered significant if $P < 0.05$. 

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Author Contributions
J.R.E., T.P.U. and T.L. supervised the project. J.R.E., S.R. and O.K.O. performed the experiments. J.R.E., S.R., A.Z.K., O.K.O., L.P. and I.G.F. analyzed the data. J.R.E., L.P., T.P.U., S.R. and D.F.C. wrote the manuscript. All authors reviewed the manuscript.

Additional Information
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