Increased rate of retinal pigment epithelial cell migration and pro-angiogenic potential ensuing from reduced cystatin C expression

Running Title: Increased RPE migration due to reduced cystatin C

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**Abstract**

**Purpose.** Variant B precursor cysteine protease inhibitor cystatin C, a known recessive risk factor for developing exudative age-related macular degeneration (AMD), presents altered intracellular trafficking and reduced secretion from retinal pigment epithelial (RPE) cells. As cystatin C inhibits multiple extracellular matrix (ECM)-degrading cathepsins, this study evaluated the role of this mutation in inducing ECM-related functional changes in RPE cellular behavior.

**Methods.** Induced pluripotent stem cells (iPSCs) gene-edited bi-allelically by CRISPR/Cas9 to express the AMD-linked cystatin C variant were differentiated to RPE cells and assayed for their ability to degrade fluorescently labelled ECM proteins. Cellular migration and adhesion on multiple ECM proteins, differences in transepithelial resistance (TER) and polarized protein secretion were tested. Vessel formation induced by gene edited cells-conditioned media was quantified using primary human dermal microvascular epithelial cells.

**Results.** Variant B cystatin C-expressing iPSC-derived RPE cells displayed a significantly higher rate of laminin and fibronectin degradation 3 days after seeding on fluorescently labelled ECM (p<0.05). Migration on matrigel, collagen IV and fibronectin was significantly faster for edited cells compared with wild-type (WT) cells. Both edited and WT cells displayed polarized secretion of cystatin C, but TER was lower in gene-edited cells after 6 weeks culture, with significantly lower expression of tight junction protein claudin-3. Media conditioned by gene-edited cells stimulated formation of significantly longer microvascular tubes (p<0.05) compared to WT-conditioned media.

**Conclusions.** Reduced levels of cystatin C lead to changes in the RPE ability to degrade, adhere and migrate supporting increased invasiveness and angiogenesis relevant for AMD pathology.

**Introduction**

The retinal pigment epithelium (RPE) comprises a unique and critical tissue in the posterior of the eye responsible for multiple aspects of maintaining visual function1. Of clinical relevance, dysfunctional and progressive loss of RPE is one of the hallmarks of late stage age-related macular degeneration (AMD)2, 3, leading to loss of central vision. This is the leading form of blindness in the elderly, but the underlying molecular mechanisms are poorly understood.

RPE cells form a tightly packed, highly polarized, monolayer between the photoreceptors and Bruch’s membrane (BrM), which in turn separates it from the choroidal blood supply. The RPE exhibits differential secretion of key effector proteins that are essential for retinal health4. One of these is the cystatin C, a 120 amino acid long protease inhibitor known to act on cysteine proteases such as cathepsins B, H and L5. The cysteine cathepsin family is comprised of 11 proteins - cathepsin B, C, F, H, K, L, O, S, V, W, and X6, which are secreted in high levels to the extracellular space where they are major players in extracellular matrix (ECM) remodelling7, 8. Chronic elevation of cathepsin activity is associated with inflammatory disease9, and can be caused by an imbalance with their endogenous inhibitors10, 11.

Multiple studies have shown that cystatin C is among the most abundantly expressed proteins in the RPE12-14, highlighting the importance of its proteolytic regulatory function in the retina. A single G73A point mutation in the cystatin C gene (CST3) has been shown to result in an increased risk of developing AMD15, 16 and Alzheimer’s disease15, 17-20, two neurodegenerative diseases that share multiple characteristics on a molecular level21. The mutation results in a variant of the protein designated variant B, which encodes an A25T amino acid substitution in the penultimate position of the highly hydrophobic secretory signal leader sequence of the precursor cystatin C. As this leader region of pre-cystatin C is cleaved off before the mature form of the protein is secreted, this mutation does not affect the function of the mature protein. However, the variant B recessive genotype has been shown to reduce the rate of cystatin C secretion22, 23, possibly as a result of intracellular mis-trafficking where an intracellular pool of mis-processed protein associates with mitochondria23.

A bottleneck in the study of RPE function has traditionally been the lack of source material, leading investigators to use spontaneously immortalized cell lines, such as ARPE19, which in some ways mimic the function of primary RPE cells24. Recent advances in the field of stem cell differentiation have led to the establishment of protocols for both spontaneous and targeted differentiation towards RPE fate25. Combined with gene editing techniques such as CRISPR/Cas9, this provides an excellent platform for studying the effect of specific protein variants in RPE cells, defining a strategy which was also employed in this study.

Our study shows that iPS-derived RPE cells gene-edited on both alleles (to reproduce the recessive disease-associated phenotype) by CRISPR/Cas9 to express the less abundant variant B form of cystatin C exhibit abnormal cellular migration and adhesion to common ECM proteins compared to wild-type (WT) cystatin C-expressing cells. The gene-edited cells also degraded fluorescently labelled laminin and fibronectin deposited into NIH3T3 fibroblast ECM 3D matrices significantly faster than non-edited cells, and although the gene-edited iPS-RPE cells formed monolayers with transepithelial resistance (TER) notably higher than the commonly used RPE cell line ARPE19, these values were significantly lower than those achieved by the non-edited cells.

**Methods**

**Cell culture**

Human induced pluripotent stem cells (iPSCs) were obtained from Human Induced Pluripotent Stem Cells Initiative (HipSci) and were cultured in a feeder-free system on 6-well plates coated with recombinant human truncated vitronectin (VTN-N; Thermo Fisher) in Essential-8 medium (E8; Thermo Fisher) at 37°C, 5% CO2. Cells were routinely passaged by dissociation with Versene (Thermo Fisher) at a 1:6 split ratio. Human ARPE19 cell line (ATCC, P25) was cultured in T75 flasks in Dulbecco’s modified Eagle’s medium/nutrient mixture F-12 Ham (DMEM:F12; Sigma) supplemented with 10% fetal bovine serum (FBS; Sigma) at 37°C, 5% CO2. Mouse NIH3T3 fibroblast cell line (ECACC) was cultured in T75 flasks in high glucose DMEM (Sigma) supplemented with 10% FBS at 37°C, 5% CO2. ARPE19 and NIH3T3 cells were routinely passaged with trypsin-EDTA solution (Sigma) and split at a 1:3-1:10 ratio. Human dermal microvascular epithelial cell (HDEMCs; PromoCell) and normal human dermal fibroblasts (NHDFs; PromoCell) were cultured in gelatin-coated T75 culture flasks in Endothelial Cell Growth Medium V2 (PromoCell) or Fibroblast Growth Medium (PromoCell), respectively, at 37°C, 5% CO2. HDMECs and NHDFs were passaged using the PromoCell DetachKit and split at a 1:3-1:5 ratio.

**CRISPR/Cas9 gene editing**

iPSCs homozygous for variant B cystatin C genotype were generated as previously described26. CRISPR/Cas9 and donor sequences (including the G73A mutation in the CST3 gene) were synthesized by System Biosciences by subcloning the AGGGATAAAACCGCAGTCGC gRNA sequence, corresponding to a section of the CST3 gene upstream of the protein coding sequence, into the CAS701R-1 EF1-T7-hspCas9-T2A-RFP-H1-gRNA plasmid, and CST3 5’- and 3’-homology arms comprising approximately 800 bp were subcloned into the HR700PA plasmid. 30 minutes before electroporation, WT iPSCs were treated with 10 µM Rock inhibitor (Y-27632, Stemgent). Cells were dissociated into single cell suspension using Stempro Accutase (Thermo Fisher) and co-transfected with the CRISPR/Cas9 and donor constructs using the Neon electroporation system (Invitrogen) with parameters 1350V, 20 ms, 1 pulse. Cells were then seeded at clonal density and cultured in E8 medium for three days (supplemented with 10 µM Rock inhibitor for the first day) followed by culture in Corning Nutristem hPSC XF medium under relevant antibiotic selective pressure for one week. Surviving clones were transferred manually to separate wells on 24 well plates using a P200 pipette and expanded in E8 medium. Genomic DNA was isolated using a DNeasy Blood & Tissue Kit (Qiagen), and successful edits on both alleles were identified via PCR and confirmed by DNA sequencing performed by DNA Sequencing & Services (MRC I PPU, School of Life Sciences, University of Dundee, Scotland, www.dnaseq.co.uk) using Applied Biosystems Big-Dye Ver 3.1 chemistry on an Applied Biosystems model 3730 automated capillary DNA sequencer. The selection cassette was then excised from the genome by LoxP-CRE recombination by transfecting the cells with CRE100A-1 plasmid (System Biosciences), seeding at clonal density and isolating single clones as described above, followed by verification through DNA sequencing. Two separate gene edited cell lines were generated from the same WT iPSC line and used in parallel throughout the study.

**Induced pluripotent stem cell differentiation to RPE**

WT or gene-edited iPSCs were passaged at least once after thawing from cryostorage and seeded on matrigel-coated (BD Biosciences) 6-well plates and grown until near confluence. Cells were differentiated towards RPE fate through a directed differentiation protocol using sequential exposure to nicotinamide and Activin A as previously described27. Pigmented patches were excised with a scalpel and seeded at a minimum of 50,000 cells per cm2. For subsequent passaging, cells were dissociated with Stempro Accutase.

**Measurement of transepithelial resistance**

100,000 iPS-RPE cells were seeded per cm2 on transwell inserts on 12-well plates and cultured for up to 8 weeks. TER was measured weekly with a Millicell ERS-2 voltohmmeter (Merck) by averaging three measurements per well and subtracting the value from a control well (containing media only) used as a baseline control. Final TER was calculated by multiplying the value with the surface area of the well (1.12 cm2).

**SDS-PAGE and western blot**

Whole cell lysates were prepared by homogenizing cells in lysis buffer (40 mM Tris, pH 6.8, 10% glycerol, 1% SDS, 5% β-mercaptoethanol, 0.01% bromophenol blue) followed by passing through a G25 needle 10 times and boiling the sample at 95°C for 5 minutes. Conditioned media from transwell experiments was collected from the upper and lower chambers after 24 hours culture and normalized to equal volume with unconditioned culture media.

Proteins were resolved by electrophoresis on 12% acrylamide gels and transferred to nitrocellulose membranes (Sigma), which were blocked with 5% milk in TBS supplemented with 0.1% Tween-20 (TBST) and probed with protein-specific antibodies. Antibodies used were against cystatin C (Abcam, ab109508), MMP2 (Cell Signaling Technology, 4022), PEDF (Millipore, MAB1059), RDH5 (Abcam, ab101457), α-tubulin (Abcam, ab4074), ZO1 (Cell Signaling Technology, 8193), ZO2 (Cell Signaling Technology, 2847), claudin-3 (Abcam, 214487), integrin α4 (Cell Signaling Technology, 8440) and integrin β1 (Cell Signaling Technology, 9699). Proteins were finally detected by ECL using SuperSignal™ West Pico PLUS Chemiluminescent Substrate (Thermo Scientific) or Radiance Plus chemiluminescent substrate (Azure Biosystems) and imaged with a ChemiDoc XRS+ imaging system (Bio-Rad).

**Pluripotency testing by immunocytochemistry**

iPSC staining of selected pluripotency markers was performed as described previously26. Briefly, iPSCs were fixed with 4% formaldehyde in PBS and probed for expression of SSEA4, OCT4, SOX2, and TRA-1-60 using a PSC 4-marker immunocytochemistry kit (Life Technologies), following manufacturer’s instructions. Immunostained cells were imaged using a Zeiss Apotome inverted fluorescence microscope.

**Cell migration assay**

iPS-RPE cells were cultured on 24-well plates coated with either matrigel, fibronectin (5 µg/cm2, Santa Cruz), laminin (5 µg/cm2, Santa Cruz), or collagen IV (5 µg/cm2, Santa Cruz) for 8 weeks. A wound approximately 500 µm wide was created by scratching the cell layer with a sterile 1000 µL pipette tip. Cells were then washed twice with DPBS to remove detached cells and cultured for up to 1 month with images of the wound recorded regularly using a Nikon Diaphot inverted light microscope. The average size of the wound was analyzed by recording 5 separate fields per well and calculating the wound area in ImageJ28, either using the MRI Wound Healing Tool (available at http://dev.mri.cnrs.fr/projects/imagej-macros/wiki/Wound\_Healing\_Tool) or by manually tracing the wound.

**Fluorescent labelling of ECM components**

100 µg of recombinant human laminin or fibronectin were labelled with a green fluorescent tag using an Alexa Fluor™ 488 microscale protein labelling kit (Thermo Fisher), following manufacturer’s instructions. Fluorescently labelled proteins were stored in the dark at -20°C as 10 µg aliquots and used within one month of preparation.

**Analysis of degradation of ECM components**

Pre-deposited ECM-matrices were prepared using previously described methods29, 30. Briefly, NIH3T3 (ECACC 93061524) mouse fibroblast cells were cultured on matrigel-coated wells on 96-well plates in the presence of 5 µg/mL fluorescently labelled laminin or fibronectin. After 1 week, cells were washed with PBS and plates were decellularised by treatment with 20 mM NH4OH (Sigma) for 5 minutes, leaving the ECM still attached to the plate, and washed twice with PBS. Plates were sealed with parafilm and stored in the dark at 4°C for up to one week before being used for downstream application.

Matrix degradation by WT or gene-edited iPS-RPE cells was analyzed by seeding 100,000 cells per cm2 on denuded fluorescent matrices and culturing for up to 72 hours with images recorded of 5 separate fields per well every 24 hours using a Zeiss Apotome inverted fluorescence microscope. Total integrated fluorescence was calculated using ImageJ software. Matrix degradation was also determined after incubation with 3-day iPS-RPE conditioned media using the same method.

**Cell adhesion assay**

iPS-derived RPE cells were dissociated into single cell suspension using Stempro Accutase and suspended into serum-free DMEM at a concentration of 5×106 cells/mL. Calcein AM (Cayman Chemical) was added to achieve a final concentration of 5 µM and the cells were incubated at 37°C for 30 min. After being washed twice in DMEM, 100,000 cells were seeded per well on a black 96-well plate pre-coated with matrigel or 5 µg/cm2 laminin, collagen IV, or fibronectin. Plates were incubated at 37°C for 1 hour, followed by washing four times with DMEM, with separate wells representing 100% adhesion remaining unwashed. Fluorescence was then measured using a Glomax-Multi+ microplate reader (Promega) equipped with a fluorescence module and blue optical kit (Ex/Em 490/510-570 nm).

**HDMEC/NHDF co-culture angiogenesis assay**

*In vitro* co-culture assay of angiogenesis was adapted from a previously described protocol31. Briefly, NHDFs (P5-10) were seeded on gelatin-coated wells on a 24-well plate at 20,000 cells per well and cultured for three days or until fully confluent. 30,000 HDMECs (P5-10) were then seeded on top of the NHDF monolayer and cultured for one day in 0.5 mL complete Endothelial Cell Growth Medium V2, after which the medium was changed to basal Endothelial Cell Growth Medium V2 supplemented with 1% FBS and 10% media conditioned by WT or CST B/B iPS-RPE cells for 24 h. In some experiments, recombinant cystatin C (Thermo Fisher) was added to the preconditioned media at a concentration of 50 ng/mL, resulting in a final co-culture concentration of 5 ng/mL. Co-cultures were maintained for five days with one media exchange, after which cells were fixed in ice cold 70% (v/v) ethanol for 30 min, blocked with 1% BSA in PBS for 30 min and immunostained against CD31 using a mouse anti-CD31 antibody (Abcam, 9498) diluted to 1 µg/mL in block buffer for 1 h. Cells were washed three times in PBS and incubated with a goat anti-mouse IgG-AP antibody (Sigma, 3562) diluted 1:500 in block buffer for 1 h. Cells were finally washed three times in distilled H2O, developed using SIGMAFAST BCIP®/NBT Alkaline Phosphatase Substrate (Sigma) and imaged using a Nikon Diaphot inverted light microscope. Five separate fields were recorded per well and analyzed using AngioQuant software32.

**Statistics**

Data are presented as means ± SD of at least three independent biological replicate experiments. Statistical analyses were performed using SigmaPlot version 13 (Systat Software, Inc). Two-tailed Student’s t-test was used to compare differences between mean values, with p-values below 0.05 considered statistically significant.

**Results**

**Gene editing and differentiation of iPSCs to RPE**

Variant B cystatin C expression has previously been studied primarily as overexpression in cell lines such as ARPE19 with reduced protein secretion being one of the notable findings23, which was also evidenced when evaluating endogenous expression in primary fibroblasts22. As the availability of cells homozygous for variant B cystatin C is low and comes with a high degree of variation between donor cells, we wanted to establish a defined platform for RPE cells homozygous for either WT or variant B cystatin C. The method for using CRISPR/Cas9 gene editing of iPSCs to generate a biallelic edit in the CST3 gene encoding an A25T mutation in the protein coding sequence has been described previously26, with a brief overview of the editing strategy shown in Figure 1A. Gene-edited iPSCs did not display any noticeable difference in morphology compared with WT cells, and to ensure that the editing process did not interfere with the pluripotent state of the cells, immunostaining of selected pluripotency markers SSEA4, OCT4, SOX2, and TRA-1-60 was performed immediately before induction of differentiation (Fig. 1B-C), with both WT and edited iPSCs displaying clear expression of all four markers. Secretion of cystatin C from gene-edited cells was confirmed to be significantly reduced compared with WT controls, while secretion of matrix metalloproteinase 2 MMP2, a protein known to be abundantly expressed by iPSCs33, was slightly elevated (Fig. 1D).

Directed differentiation of iPSCs into RPE-like cells was achieved by employing a previously described protocol27, with pigmented patches starting to appear after approximately one month of culture. At this stage, it became apparent that the pigmentation of gene-edited cells was not as strong as seen in WT cells (Fig. 2A). While WT cells displayed defined black spots of hyperpigmented cells, the spots seen in edited cell cultures were fainter with diffuse borders. Patches were manually excised and subcultured on newly prepared matrigel-coated plates and after approximately one month of further culture WT and edited iPS-RPE cells had both established tight monolayers of cells. Only WT cells displayed sections with the cobblestone phenotype characteristic for RPE (Fig. 2A), and pigmentation was noticeably stronger in WT cells, phenotypes which continued to be observed during following passages. Expression of visual cycle protein and RPE marker RDH5 was confirmed by Western blot analysis for both WT and edited cells, as well as secretion of PEDF (Fig. 2B). Two individual gene-edited clones were differentiated separately and analyzed by functional assays throughout this study.

**Effect of reduced cystatin C expression on monolayer polarity**

To evaluate whether editing of cystatin C gene had any effect on polarized secretion and TER, cells were cultured on transwell inserts allowing for separate collection of media conditioned to the apical or basolateral side of the monolayer. The human ARPE19 spontaneously immortalized RPE cell line was cultured in a similar manner and used for comparison. As seen in Figure 3A, ARPE19 TER quickly reached its highest point of approximately 50 Ω × cm2 after just one week in culture, a value consistent with previous studies24, 34, while the iPS-derived RPE cells slowly matured towards significantly higher levels. CST3 B/B cells reached their maximum levels of approximately 150-200 Ω × cm2 after 4 weeks in culture, while non-edited cells displayed maximum values approaching 300 Ω × cm2 after 6 weeks in culture. Although these levels are lower than those observed for *in vitro* cultures of primary RPE cells34-36, they are a clear improvement over the ARPE19 model cell line, prompting us to investigate whether any polarization of secreted proteins could be observed. Western blot analysis of proteins secreted from the apical and basolateral side of the monolayer showed a clear polarization of cystatin C secretion from both WT and CST3 B/B cells (Fig. 3B). To demonstrate that this predominant basolateral secretion was not a global feature of the cells, we also probed the membranes against PEDF, a protein previously shown to be predominantly secreted apically from RPE cells, where the abundance was not significantly different between the two sides (Fig. 3B). TER is known to be a function of tight junction strands37 comprised in RPE of proteins such as ZO1, ZO2, claudins and occludin, prompting us to investigate whether CST3 B/B cells displayed reduced expression of these proteins. Western blot analysis of iPS-RPE whole cell lysates showed no significant difference in expression of tight junction proteins ZO1 or ZO2; however, claudin-3 expression was significantly reduced in CST3 B/B cells (Fig. 3C-D).

**Effect of reduced cystatin C expression on ECM composition**

Cystatin C is one of the most abundantly expressed proteins in the RPE13 and mainly acts as an inhibitor of various proteases, many of which are known to cleave commonly expressed ECM proteins such as laminin and fibronectin7. To evaluate whether the reduced expression levels of cystatin C in our gene-edited iPS-RPE cells were associated with a higher rate of ECM degradation, we first incorporated fluorescently labelled laminin and fibronectin into ECM deposited onto matrigel-coated plates by mouse fibroblast cells. Cells were then lysed via NH4OH exposure, leaving the deposited ECM intact. iPS-RPE cells were seeded on top and cultured for up to 3 days with daily monitoring by fluorescence microscopy. As seen in Figure 4A-B, gene-edited iPS-RPE cells were able to degrade both laminin and fibronectin in the deposited ECM significantly faster than the WT cells. As cystatin C is a potent inhibitor of secreted proteases, we suspected that this effect may have been due to the loss of cystatin C regulatory effect in the RPE secretome. However, incubation of fluorescent ECM matrix with media conditioned to either WT or gene-edited iPS-RPE cells for up to 3 days yielded no discernible difference in rate of degradation (Fig. 4B). From these results, it became clear that the increased rate of laminin and fibronectin degradation required the physical contact to the RPE cells.

**Effect on cell adherence and migration on ECM**

To investigate if cellular adhesion and migration are affected in gene-edited cells expressing variant B cystatin C, tissue culture plates were coated with common ECM proteins laminin, collagen IV and fibronectin, , as well as matrigel, which is comprised by a complex mixture of proteins. iPS-RPE cells loaded with the fluorescent dye calcein AM were seeded onto these plates, followed by washing after one hour of allowing the cells to attach to the well surface. Gene-edited cells adhered to the plates more efficiently than WT cells when seeded on collagen IV and fibronectin (Fig. 5A). The difference in rate of adherence to matrigel and laminin were not statistically significant between WT and edited iPS-RPE. Cellular adherence to ECM is largely dependent on interactions with integrin receptors, and we detected a notable increase in expression of integrin α4 and β1 in edited iPS-RPE cells (Fig. 5B).

The ability of cells to migrate on the different surfaces was also tested by allowing the cells to establish confluent monolayers on the different coatings and inducing a wound using a sterile pipette tip. Frequent monitoring by light microscopy revealed that gene-edited cells migrated significantly faster on all tested surfaces except laminin-coated plates (Fig. 6). The difference was especially striking when cultured on matrigel for 7 days, at which edited cells had achieved near complete closure while WT cells had barely begun to close the wound.

**Reduced cystatin C in gene edited iPS-RPE-conditioned media supports a pro-angiogenic environment**

Media conditioned by WT or gene edited iPS-RPE cells collected after 24h was supplemented to the culture media used in a NHDF/HDMEC co-culture angiogenesis assay to investigate whether the changes in secretion seen in the edited cells were better at promoting microvascular tube formation. Indeed, we found that co-cultures using the media conditioned by cystatin C gene edited iPS-RPE cells achieved a clear difference in HDMEC tubular morphology, with significantly longer tubules formed (Fig. 7). This effect could be offset by adding exogenous cystatin C to the media at a concentration of 50 ng/mL, which is approximately the concentration we previously detected in confluent cultures of primary fetal RPE cells after 24h36.

**Discussion**

The gene-edited iPS-RPE cell lines presented reduced cystatin C secretion, consistent with previous studies on variant B cystatin C genotype in cultured cells22, 23, 38, serum plasma39, or cerebrospinal fluid40, 41, with one exception - a study by Nguyen et al. that did not observe a functional effect on secretion rate conferred by the mutation when overexpressed in HEK293T and non-differentiated ARPE19 cells42. The present study also noted significantly lower levels of intracellular cystatin C in gene-edited cells, highlighting that expression level was affected. An increased risk for developing AMD, as well as other degenerative diseases such as Alzheimer’s disease15, 17-20 has been attributed to this mutation, making functional studies on how the cells behave in defined conditions valuable. At the same time, the Beaver Dam Eye Study identified a correlation between increased cystatin C serum levels and AMD pathology43, further complicating the functional relationship between cystatin C and AMD. Elevated cystatin C serum concentration was also described as a risk factor for other disease conditions, e.g. cardiovascular disease44, 45 and progression to pre-diabetes46. However, significantly reduced serum cystatin C levels, which had been correlated with variant haplotypes47, 48, were highlighted in association with atherosclerotic lesions, dilated aorta and focal progression of coronary artery disease47, 49, 50. These data on involvement of cystatin C proteostasis in cardiovascular disease suggested that cystatin C deficiency alters arterial structures through impaired regulation of proteolysis of the arterial ECM.

Both WT and edited iPSCs could be differentiated to RPE phenotype using a previously described targeted differentiation protocol27. We favored a targeted instead of spontaneous differentiation protocol mainly due to being able to maintain consistency between experiments. Hyperpigmented patches of RPE-like cells appeared at identical time points for both WT and edited cells, which could be manually excised and further subcultured for multiple passages. Clear differences in morphology were observed between WT and CST3 B/B iPS-RPE cells, although immunoblot analysis of whole cell lysates showed no significant difference in expression of visual cycle and tight junction proteins. Gene-edited cells appeared smaller and less pigmented after differentiation, and adopted less regular hexagonal patterns, which could point to RPE developmental problems.

The overall process for obtaining genetically defined RPE cells described here could easily be applied towards functional studies of other genotypes associated with retinal diseases, and as the gene editing is done at the iPSC stage, the cells could also hypothetically be differentiated to other relevant cell types.

The level of reduction in cystatin C secretion seen in our gene-edited cells was dramatic, but consistent between two separate cell lines generated. A previous study in primary CST3 A/A, A/B and B/B skin fibroblast cells demonstrated high level of variation in cystatin C secretion between donors of the same genotype22, highlighting the value of using non-edited cells, which are otherwise genetically the same, as controls. However, in their study the expression levels seen in whole cell lysates were not significantly affected by the B/B genotype, while we observed a clear reduction in expression levels. It is possible that this effect is cell-type dependent, or that the remaining scar sequence from gene editing located upstream of exon 1 (Fig. 1A) affects the level of protein expression, as a previous study has identified this region to be sensitive to modifications51, thus accentuating the reduction of secretion.

As cystatin C is a main regulator for multiple cysteine proteases known to degrade ECM proteins, the effects of reduced protein expression and secretion have on cellular interaction with ECM structures was our main focus in this study. Cystatin C is predominantly secreted from the basolateral side of the RPE monolayer, which is located directly adjacent to the collagen-rich BrM, and is therefore likely to in part regulate BrM maintenance and remodeling. With both age and AMD progression, the BrM is known to experience thickening52, a likely result of progressive loss of proteostatic control. Whether cystatin C plays a role in this process is yet to be established.

Changes in migration and adhesion of gene-edited iPS-RPE cells appear to be substrate dependent, as neither was significantly altered compared with WT cells when cultured on laminin-coated wells. This suggests active mechanistic functions being affected, rather than general cellular properties. As laminin is the predominant component of matrigel (approx. 60% (w/w)), we were surprised to see that the migration rate on matrigel was dramatically increased in edited iPS-RPE cells, while no significant difference in adhesion compared with WT cells was seen. Experiments conducted on collagen IV, which is the second most abundant component of matrigel (approx. 30% (w/w)), resulted in clear alteration of both adhesion and migration in edited cells.

Analyzing whole cell lysates, we detected a clear increase in expression of multiple integrins. These transmembrane receptor proteins form heterodimers in different combinations that provide interactions between the intracellular cytoskeleton and extracellular ECM molecules, thereby playing key roles in cellular adhesion53. Specifically, we detected marked increases in integrins α4 and β1 expression. The integrin α4β1 heterodimer recognizes fibronectin54, to which edited iPS-RPE cells displayed increased adhesion, although both chains are also involved in other combinations that mediate interaction with laminins and collagens. It is possible that integrin expression becomes upregulated in response to increased degradation rate of ECM coatings on the tissue culture wells, which would explain the increased rate of adhesion to fresh coatings when subcultured.

Average content of tight junction protein claudin-3 was significantly reduced in our gene edited iPS-RPE cells. This is the second most abundantly expressed claudin in RPE cells55, and crucially, depletion of claudins has been shown to have the potential to drastically disrupt TER in cultured RPE cells55. This may in part explain why our edited iPS-RPE cells also display a significant reduction of TER after 6 weeks in culture.

To functionally test the specific link between our model and exudative AMD phenotype, we investigated how changes in cystatin C expression and secretion affect microvascular tube formation in an NHDF/HDMEC co-culture angiogenesis model. The dysregulated secretion seen in our gene-edited cells resulted in an increased ability for HDMECs to form vessels when cultured in media preconditioned by the variant B cystatin C iPS-RPE cells. As the effect was eliminated when supplementing the media with exogenous cystatin C, this demonstrates a functional association between cystatin C and angiogenesis, further underlining the clinical relevance of cystatin C as an effector molecule in the context of wet AMD.

In conclusion, reduced expression of cystatin C in iPS-derived RPE cells had clear effects on cells’ behavior and interaction with ECM. In this study, reduced expression was induced via CRISPR/Cas9 gene editing, but reduced protein levels have previously been described in human fetal RPE cells cultured on ECM subjected to advanced glycation end products and a general decline with age in posterior RPE segments36. The evidence presented provides a functional link between reduced cystatin C expression and RPE invasiveness in a pro-angiogenic environment relevant for wet AMD pathology, affording mechanistic data explaining the functional relationship between cystatin C and cellular ECM interactions.

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**Figure legends**

**Figure 1. Bi-allelic gene editing of iPSCs to encode variant B cystatin C.**

(A) Overall schematic of strategy used for gene editing of iPSCs (top). Sequencing of the CST3 gene confirmed a homozygous G73A edit (bottom). Following gene editing, pluripotency staining of WT (B) and edited (C) cells using antibodies against SOX2, TRA-1-60, SSEA4 and OCT4 demonstrated that the editing had not affected the pluripotent state of the cells. Scale bars, 200 µm. (D) Western blot analysis of conditioned media from two separate clones showed that cystatin C secretion was strongly reduced. Images shown are representative of 3 individual experiments.

**Figure 2. Differentiation of gene-edited iPSCs to RPE cells.**

(A) Hyperpigmented patches of RPE cells following differentiation were excised at P0 (upper panels) and further expanded. Lower panels show cells at P2. Arrows, pigmented patches prior to excision during P0; Scale bars 250 µm. (B) Western blot analysis of conditioned media and whole cells lysates confirmed reduced cystatin C expression and secretion, as well as expression of RDH5 and secretion of PEDF. Images shown are representative of 3 individual experiments.

**Figure 3. Reduced TER in CST3 B/B iPS-RPE cells.**

(A) TER of WT and gene-edited iPS-RPE cells seeded on transwell inserts was monitored over 8 weeks. (B) Conditioned media from the upper and lower wells of transwell inserts, corresponding to apical and basolateral secretion, were analyzed by Western blot for cystatin C and PEDF secretion. A, apical secretion; B, basolateral secretion. (C) Expression of tight junction proteins ZO1, ZO2 and claudin 3 was confirmed by Western blot. Images shown are representative of 3 individual experiments. (D) Densitometric analysis of tight junction protein content from whole cell lysates shown in panel C. n = 3; \*, P ≤ 0.05.

**Figure 4. Faster degradation of ECM by CST3 B/B iPS-RPE cells.**

(A) Degradation of fluorescently labelled laminin and fibronectin incorporated into NIH3T3 fibroblast ECM 3D-matrix by iPS-RPE cells or conditioned media was analyzed by fluorescence microscopy. NT, no treatment; Scale bars 200 µm. (B) Quantitative image analysis of total integrated fluorescence. Gene-edited iPS-RPE cells were able to degrade both laminin and fibronectin faster than WT cells, while no difference in degradation rate could be seen when subjecting the labelled ECM to conditioned media.

**Figure 5. Increased adhesion to ECM coated wells by CST3 B/B iPS-RPE cells.**

(A) iPS-RPE cells were subjected to cell adhesion assays on wells coated with matrigel, laminin, collagen IV, or fibronectin. A significantly larger proportion of gene-edited cells were able to attach to wells coated with collagen IV and fibronectin, compared with WT cells, after 1 hour. No significant difference could be seen when seeding on wells coated with matrigel or laminin. n = 3; \*, P ≤ 0.05. (B) Western blot analysis of iPS-RPE whole cell lysates shows increased expression of integrin α4 and β1 in CST3 B/B cells. Images shown are representative of 3 individual experiments.

**Figure 6. Faster wound closure of CST3 B/B iPS-RPE cells.**

Wound healing assays were conducted with iPS-RPE cells cultured on matrigel, laminin, collagen IV, or fibronectin. Wound closure of gene-edited iPS-RPE cells was significantly higher after 7 days when cultured on matrigel, collagen IV and fibronectin. No significant difference could be seen between WT and gene edited iPS-RPE cells when cultured on laminin. n = 3; \*, P ≤ 0.05.

**Figure 7. Increased angiogenesis supported by media conditioned by CST3 B/B iPS-RPE cells.**

(A) NHDF/HDMEC co-cultures were stimulated with media conditioned by WT or CST3 B/B iPS-RPE cells, followed by microvascular staining with anti-CD31 antibody and analysis by light microscopy. Images shown are representative of fields from three separate experiments. (B) Quantitative analysis of microvascular networks from NHDF/HDMEC co-culture experiments. n = 3; \*, P ≤ 0.05.