Prominin-1 Is a Novel Regulator of Autophagy in the Human Retinal Pigment Epithelium

Sujoy Bhattacharya, Jinggang Yin, Christina S. Winborn, Qiuhua Zhang, Junming Yue, and Edward Chaum

1Department of Ophthalmology, University of Tennessee Health Science Center, Memphis, Tennessee, United States
2Department of Pathology, University of Tennessee Health Science Center, Memphis, Tennessee, United States
3Department of Anatomy and Neurobiology, University of Tennessee Health Science Center, Memphis, Tennessee, United States

Correspondence: Edward Chaum, Plough Foundation Professor of Retinal Diseases, UTHSC Hamilton Eye Institute, 930 Madison Avenue, Memphis, TN 38163, USA; echaum@uthsc.edu.
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PURPOSE. Prominin-1 (Prom1) is a transmembrane glycoprotein, which is expressed in stem cell lineages, and has recently been implicated in cancer stem cell survival. Mutations in the Prom1 gene have been shown to disrupt photoreceptor disk morphogenesis and cause an autosomal dominant form of Stargardt-like macular dystrophy (STGD4). Despite the apparent structural role of Prom1 in photoreceptors, its role in other cells of the retina is unknown. The purpose of this study is to investigate the role of Prom1 in the highly metabolically active cells of the retinal pigment epithelium (RPE).

METHODS. Lentiviral siRNA and the genome editing CRISPR/Cas9 system were used to knock out Prom1 in primary RPE and ARPE-19 cells, respectively. Western blotting, confocal microscopy, and flow sight imaging cytometry assays were used to quantify autophagy flux. Immunoprecipitation was used to detect Prom1 interacting proteins.

RESULTS. Our studies demonstrate that Prom1 is primarily a cytosolic protein in the RPE. Stress signals and physiological aging robustly increase autophagy with concomitant upregulation of Prom1 expression. Knockout of Prom1 increased mTORC1 and mTORC2 signaling, decreased autophagosome trafficking to the lysosome, increased p62 accumulation, and inhibited autophagic puncta induced by activators of autophagy. Conversely, ectopic overexpression of Prom1 inhibited mTORC1 and mTORC2 activities, and potentiated autophagy flux. Through interactions with p62 and HDAC6, Prom1 regulates autophagosome maturation and trafficking, suggesting a new cytoplasmic role of Prom1 in RPE function.

CONCLUSIONS. Our results demonstrate that Prom1 plays a key role in the regulation of autophagy via upstream suppression of mTOR signaling and also acting as a component of a macromolecular scaffold involving p62 and HDAC6.

Keywords: CD133, Atg5, Atg7, LC3-II, Akt, p62, mTORC1, mTORC2, autophagy flux, Nutlin-3
Autophagy is a highly conserved cellular process that plays a central role in metabolism, proliferation, and survival. It is involved in the degradation of cellular components, including organelles, proteins, and macromolecules. Autophagy can be triggered by various stimuli, including nutrient deprivation, cellular stress, and aging, and it serves as a protective mechanism against cellular degradation and death.

**Methods**

**Reagents**

Disposable cell culture ware was purchased commercially (Corning Glass Works, Corning, NY, USA). The ARPE-19 cell line was purchased from ATCC CRL-2302 (Manassas, VA, USA), and the human retinal endothelial cell (HREC) line was a kind gift from Jena Steinle. Cell culture medium was obtained from Lonza (Walkersville, MD, USA), and fetal bovine serum (FBS) was purchased from Gibco (Life Technologies, Carlsbad, CA, USA). All chemicals were of the highest purity commercially available.

Lentiviral vectors containing human Prom1 siRNA (pLenti-Prom1-siRNA-GFP) were purchased from Applied Biological Materials, Inc. (Richmond, Canada). A dual convergent promoter system where two different promoters (U6 and H1 promoters) controlled the sense and antisense strands of the siRNA was used. Four siRNA lentiviral constructs (Prom1-603-siRNA, Prom1-736-siRNA, Prom1-1087-siRNA, Prom1-1270-siRNA) were used to knock down Prom1 expression in RPE cells. Four constructs were transfected in primary RPE cells using Fugene-6 HD. GFP expression by live cell confocal microscopy was used to monitor transfection efficiency. Forty-eight hours posttransfection, RPE cells were treated with puromycin to enrich cultures expressing Prom1 siRNA. GFP-positive stable transfectants with puromycin resistance were used for further analyses. Western blotting with Prom1 antibody was used to evaluate Prom1 expression in cells stably transfected with lentiviral-siRNA.

**Confocal Microscopy**

Confluent RPE and HREC cell monolayers were fixed in ice-cold Acetone/Methanol (1:1) followed by permeabilization in 0.1% Triton X-100. After permeabilization, the monolayers were blocked in 5% BSA in PBS and further incubated with primary antibodies (rabbit polyclonal Prom1, mouse β-catenin, rabbit LC3-I and LC3-II) for 1 hour at 37°C, followed by 1 hour incubation with secondary antibodies (Alexa Fluor 488-conjugated anti-mouse IgG and Cy3-conjugated anti-rabbit IgG antibodies). For LC3 staining, cells were mounted on glass slides using mounting medium containing 4',6-diamidino-2-phenylindole (DAPI). For localization of Prom1 and β-catenin, the fluorescence was examined under a Zeiss LSM 5 laser scanner confocal microscope, and images were collected using LSM 5 Pascal software as described earlier. Images were stacked using the software, Image J (National Institutes of Health, Bethesda, MD, USA), and processed by Adobe Photoshop (Adobe Systems, Inc., San Jose, CA, USA).

**Prom1 Lentiviral Construct**

The wild-type (WT) Prom1 lentiviral construct (pLenti-GII-CMV-GFP-2A-Puro; NM_001145848.1) was obtained from Applied Biological Materials, Inc., expressing Prom1 under the control of CMV promoter. Plasmids were amplified using the Qiagen Plasmid maxi prep kit following the manufacturer’s instructions. Ten micrograms Prom1 plasmid and the empty GFP control plasmid were packaged in human 293FT cells and purified through ultracentrifugation at the Viral Vector Core laboratory as described previously.
CRISPR-Cas9-Mediated Genomic Deletion of Prom1

The 17-nucleotide guide RNA (gRNA) sequence (5'-GGATGCACCAAGGACAGCAG-3') was used to target human PROM1 gene at exon 10. Oligonucleotides were purchased from Integrated DNA technology and annealed, followed by phosphorylation through T4 DNA polynucleotide kinase reaction. The annealed double-stranded DNA was cloned into the BsmBI-BsmBI sites downstream from the human U6 promoter in the Lent-CRISPR v2 plasmid (Addgene plasmid #52961). The Lent-CRISPR-Prom1 construct was packaged into 293FT cells and purified as described previously. Purified lentivirus was used to infect ARPE-19 cells. During infection, 15 µL of the purified lentivirus (Cas9 or Prom1-Cas9) was added to 60% to 70% confluent ARPE-19 cells. During infection, 15 µL of the purified lentivirus (Cas9 or Prom1-Cas9) was added to 60% to 70% confluent ARPE-19 cells. 

Overexpression and KO of Prom1 were verified by genomic DNA analysis. Control ARPE-19 cells, and cells infected with empty Cas9, WT Prom1, and CRISPR Cas9 lentivirus were used to extract genomic DNA using the Qiagen micro kit. PCR was performed using the forward primer: 5'-TATGGTACCAATACTGAAGAAG-3' and reverse primer: 5'-TAGTTGGAGCAGCTGTTAGAGCA-3'. The single band PCR product was confirmed using a 1.2% agarose gel for each cell line. The PCR products were purified with the QIA quick PCR Purification kit and quantified by Nano Drop Spectrophotometry. The purified PCR products were inserted into the pCR2.1 vector using the TA Cloning kit (Invitrogen). For each try, 100 ng PCR product, 12.5 ng pCR2.1 vector, 2.5 U vector using the TA Cloning kit (Invitrogen).PCR was performed using the forward primer: 5'-GATGACCGCAGGCT-3' and reverse: 5'-AGACGCTGAGTTACATTG-3'. The primer sequences were used. The double-stranded DNA was cloned into the BsmBI–BsmBI sites through T4 DNA polynucleotide kinase reaction. The annealed DNA was ligated and the total number of LC3-1/II were identified by incubation with 1 µM TO-PRO3 (Invitrogen). Flow cytometry, control and Prom1-KO cells were treated as described above, rinsed with PBS, and trypsinized. Cells were collected by centrifugation, fixed in ice-cold methanol for 20 to 30 minutes at –20°C or 2% paraformaldehyde for 15 minutes at 25°C, and permeabilized in PERM buffer (BD Biosciences, San Jose, CA, USA) for 30 minutes on ice.
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followed by blocking with FcγR (BioLegend, San Diego, CA, USA) for 10 minutes. Cells were incubated with Prom1 (1:100), LC3-I/II (1:100) or 0.5 μg LAMP2 (Abcam, Cambridge, MA, USA) antibodies for 1 hour at room temperature, washed, and incubated with Alexa Fluor 488 and Alexa Fluor 647 secondary antibodies (Thermo Fisher Scientific) for 45 minutes. Samples were washed and analyzed with the FlowSight Imaging Flow Cytometer (EMD Biosciences/Millipore Corp.), which simultaneously produced dark field (side scatter), bright field (BF), and fluorescence images at ~20× magnification. Compensation controls were obtained from single color stained cells. The IDEAS software (Amnis Technology, EMD Millipore, Billerica, MA, USA) was used to separate single cells and cell doublets positive for CD133+/LAMP2+ or LC3+/LAMP2+ cells. We used the Spot Counting Wizard in IDEAS to automatically quantify LC3 puncta formation. In order for the Wizard to correctly identify LC3 puncta staining, we manually identified cells expressing low (diffuse) and high (punctate LC3) numbers of LC3 puncta. The wizard used these populations to generate a spot mask, which was applied to count individual spots across our sample population (5000–10,000 cells). A spot count of 3 or greater was considered punctate. Cells with variable levels of LC3 intensity were chosen for the reference population to avoid bias based on LC3 intensity. LC3-I and LC3-II protein levels were further analyzed by western blotting. To quantify autophagy in RPE cells, we compared (1) the levels of LC3-II to LC3-I and (2) levels of LC3-II to actin. Both quantification methods demonstrated similar conclusions throughout our studies involving primary RPE and ARPE-19 cells.

Relative Hypoxia

Primary RPE cells seeded in six-well clusters were cultured at 20% O₂ and 5% CO₂. After reaching 60% to 70% confluency, the culture medium was changed and treated as 0 hours. For the relative hypoxia experiments, cells were subsequently incubated at 37°C in a controlled environment of 5% CO₂, 8% O₂, and 87% N₂ for the specified time periods. Cells cultured under hypoxic conditions were immediately processed and stored at ~80°C.

STR Analysis

The batch of the ARPE-19 cells used in this study was validated by short tandem repeat (STR) analysis performed by the ATCC Cell line authentication service. Briefly, 17 STR loci plus the gender-determining locus, Amelogenin, were amplified using the commercially available PowerPlex 18D kit from Promega, and the cell line sample was processed using the ABI Prism 3500xl Genetic Analyzer. Data were analyzed using GeneMapper ID-X v1.2 software (Applied Biosystems). Our cells were found to be a perfect match for the ATCC human RPE cell line CRL-2302 (ARPE-19). The STR analysis result is a Supplementary File.

Primary RPE and HREC Cultures

Primary human RPE cells were isolated from postmortem deidentified donor eyes provided by the Mid South Eye Bank. The Institutional Review Board at the University of Tennessee Health Science Center approved the use of human eyes from deidentified donors. We used primary RPE cells that were derived from two young donors (age 29 or 40) and from two aged donors (>70 years of age) as described previously.32 All donor eyes were shipped to our laboratory within 24 hours of enucleation. Globes were excised, anterior segment was removed, vitreous was extracted manually, and the retina was dissected free. The eyecup was washed three times with Dulbecco's modified Eagle's medium (DMEM), and 0.25% trypsin/EDTA was applied for four 15-minute digestion cycles. Cells were loosened by aspiration, transferred to DMEM with FBS, spun at 2000g for 5 minutes, and the cell pellet was reseeded in DMEM in 15% FBS and plated in poly-L-Lysine coated 12-well cell culture ware. The fastest growing cells with cobblestone morphology were used for our studies. Primary cultures within the first three to five passages were used for our studies. Stock cells were maintained in DMEM and Ham’s F12 medium (1:1) ratio containing L-glutamine and 10% FBS in a humidified, 37°C incubator in an atmosphere of 5% CO₂. RPE cells were cultured using protocols described previously.33 Briefly, RPE cells were seeded on plastic cell wares and confluent monolayers were used for experiments. For differentiating cultures, RPE cells were seeded on transwell inserts, and the cells were grown for more than 4 weeks in DMEM containing 1% FBS. The HRECs were cultured in cell-ware pretreated with attachment factor in DMEM:F12 (1:1) media containing 1% penicillin-streptomycin, endothelial cell growth supplement (ECGS; Sigma-Aldrich Corp.) and 10% FBS and grown in 5% CO₂ at 37°C. Medium was changed every 2 days, and cells between three and five passages were used for all experiments.

Western Blotting

Cell lysates were prepared using mammalian protein extraction buffer (Pierce, Rockford, IL, USA) with 150 mM NaCl, 1 mM Na₂ EDTA and a protease inhibitor cocktail followed by SDS-PAGE. Proteins were transferred to Immobilon-P membranes (Millipore, Bedford, MA, USA) and probed with primary antibodies overnight at 4°C in Tris-buffered saline (TBS) containing 0.1% Tween-20 and 5% nonfat dry milk (Bio-rad, Hercules, CA, USA). Membranes were subsequently incubated with horseradish peroxidase-conjugated secondary antibodies at room temperature for 1 hour, and the immunocomplexes were visualized by the ECL detection system (Perkin Elmer, Hercules, CA, USA) using the Kodak Image Station 4000R. Membranes were stripped and reprobed for actin or GAPDH as loading controls. Representative western blots from three experiments are shown. Densitometric analysis of all western blots was performed using Image J software (developed by Wayne Rasband, available at http://rsb.info.nih.gov/ij/index.html, provided in the public domain by the National Institutes of Health).

Immunoprecipitation

RPE cells were rinsed with ice cold PBS and lysed by freeze thawing in NP40 cell lysis buffer (Invitrogen) containing protease and phosphatase inhibitors (Thermo Fisher Scientific). The lysates were transferred to Eppendorf tubes and centrifuged at 12,500 rpm for 15 minutes at 4°C. The cell extracts containing equal amounts of proteins were incubated with the appropriate antibodies overnight at 4°C, followed by the addition of protein A/B Sepharose CL4B beads (GE Healthcare 71-7089-00 AE) with gentle rocking for 2 hours. The beads were washed three times with lysis buffer and once with PBS, and the immunocomplexes were released by heating in Laemmli sample buffer and analyzed by western blotting using specific antibodies.

Statistical Analysis

All data are expressed as mean ± SE. Experiments were repeated three times, with triplicate samples for each. ANOVA and Bonferroni post hoc testing determined the significance of
the differences between means. Values of $P < 0.05$ were considered significant.

RESULTS

Expression and Localization of Prom1 in RPE Cells

We investigated the expression and localization of Prom1 in both immortalized ARPE-19 cells and primary RPE cultures obtained from donor eyes. Similar levels of Prom1 expression were observed in ARPE-19 cells and primary RPE cultures (Fig. 1A). Immunofluorescence staining of Prom1 and β-catenin expression show that Prom1 is mainly distributed in the cytoplasm and perinuclear regions with some nuclear staining in the ARPE-19 and RPE cultures, which failed to co-localize with peripheral β-catenin (Fig. 1B). Prom1 does co-localize with β-catenin in HRECs, confirming that the antibody can detect Prom1 in the cell membrane. This suggests that, surprisingly, Prom1 is intracellular and does not localize to the RPE cell membranes at the intercellular junctions.

To further investigate Prom1 expression, we used confluent primary RPE and ARPE-19 monolayers grown on plastic cell culture-wares or differentiated RPE cultures using prolonged growth on transwell inserts as performed previously, and examined Prom1 expression using western blotting. Differentiation of RPE was associated with the increased expression of ZO-1 and β-catenin and decrease in Prom1 expression, compared to nondifferentiated RPE (Fig. 1C), suggesting a correlation between reduced Prom1 expression and RPE differentiation.

Prom1 Regulates Autophagy in RPE Cells

Since Prom1 localized to the cytoplasm in RPE cells (Fig. 1B) and cytoplasmic localization of Prom1 has been correlated with increased autophagy in hepatoma cells, we performed experiments to investigate the role of Prom1 in autophagy in the RPE. Prom1 expression was significantly downregulated in
RPE cells stably transfected with Prom1-siRNA (construct 1087, Fig. 2A) and partially blocked when all four siRNA constructs were used. Upon induction of autophagy, microtubule-associated protein light chain-I (LC3-I) is conjugated by Atg7, Atg3, and Atg-5 multimers to the lipophilic phosphatidyl-ethanolamine to generate LC3-phosphatidylethanolamine conjugate (LC3-II), which is recruited to autophagosomal membranes and whose levels correlate with autophagosome number. 35,36 Knockdown of Prom1 with construct 1087 dramatically decreased levels of both LC3-I and LC3-II (Fig. 2A), suggesting that Prom1 is a novel regulator of autophagy in RPE cells. The mixture of all Prom1-siRNA constructs reduced Prom1, LC3-I, and completely reduced LC3-II expression. Since autophagy has been implicated in RPE survival, we hypothesized that inhibition of basal RPE autophagy by Prom1 siRNA might be detrimental for RPE homeostasis, perhaps leading to the induction of caspase-3-dependent apoptosis. Therefore, we measured caspase-3 activation in cells transfected with Prom1 siRNA, and found that caspase-3 was not activated in these cells. These results indicate that inhibition of basal RPE autophagy failed to induce RPE apoptosis (Fig. 2A).

### Prom1 Expression Correlates With the Induction of Autophagy in Response to Stressors and Aging in RPE Cells

Extracellular and intracellular stress signals, like hypoxia, oxidative stress, and nutrient deprivation, are known to trigger autophagy. We next examined the role of Prom1 expression on hypoxia- or nutrient deprivation-induced...
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Acute hypoxic exposure (1%-3% oxygen) of RPE cells increases expression of both hypoxic master regulator, HIF-1α, and angiogenic stimulator, VEGF within 12 hours.38,39 Since reduced oxygen ~12% is physiologic and is sufficient to induce moderate hypoxia in humans,40,41 we exposed RPE cells to moderate hypoxia (1% oxygen) for 0 to 96 hours and examined the effect of hypoxia on autophagy and Prom1 expression. HIF-1α expression significantly increased after prolonged hypoxic treatment of 72 hours and remained elevated at 96 hours (Fig. 2B). Autophagy is a dynamic process where LC3-I is rapidly converted to LC3-II. However, the pattern of LC3-I (precursor) to LC3-II (product) conversion is not only cell type specific, but also related to the type of stimulus and the relative levels of LC3-I and LC3-II proteins in various cell types.42 Primary RPE cells exposed to hypoxia for 24 hours showed a significant reduction of LC3-I without increasing LC3-II levels (Fig. 2B). Both LC3-I and LC3-II levels showed a similar pattern after 48 hours of hypoxia, indicating a rapid interconversion of LC3-I to LC3-II followed by consumption of the LC3-II protein. Both LC3-I and LC3-II protein levels increased after 72 hours of relative hypoxia. However, LC3-I levels decreased and LC3-II expression remained elevated after 96 hours of hypoxia. Although LC3-II levels are commonly normalized to actin to measure autophagy, actin levels may decrease when autophagy is induced in many organisms ranging from yeast to mammals.42 Furthermore, ignoring the changes in LC3-I in favor of LC3-II normalized with actin13 may not provide an accurate picture of the autophagic response in RPE cells. Since the ratio of LC3-II to LC3-I was previously used to monitor autophagy flux in the retina,44 we used LC3-II/LC3-I ratio as an index for autophagy activation in primary RPE cells. Measuring LC3-II to LC3-I ratio showed activation of autophagy in primary RPE cells within 24 hours of moderate hypoxia, which increased throughout the entire time period of hypoxia treatment (Fig. 2C). BAF is a specific inhibitor of vacuolar-type ATPase (V-ATPase) and is known to prevent the fusion of autophagosomes with lysosomes resulting in an inhibition of autophagy.45 Exposure of primary RPE cells with 48 hours of hypoxia in the presence of BAF further increased LC3-I and LC3-II accumulation, suggesting activation of autophagy flux in response to hypoxia (Supplementary Fig. S1). To confirm whether cells exposed to 96 hours of hypoxia were undergoing apoptosis, samples were analyzed for activation of caspase-3. Data presented demonstrate that moderate hypoxia for 96 hours failed to activate caspase-3, indicating sustained autophagy exerts a protective effect during hypoxia in primary RPE cells. p62 is also commonly used as a marker for the induction of autophagic flux46 as p62 levels are inversely correlated with the induction of autophagy. Moderate hypoxia reduced p62 expression at 24 hours, which continued to decline in a time-dependent manner (Figs. 2B, 2C). Likewise, phosphorylation of p62 Ser349 decreased in response to hypoxia in a time-dependent manner. Exposure to hypoxia for 72 to 96 hours significantly increased expression of Prom1, Atg5, and LC3-II/LC3-I ratio, without altering Atg3 expression (Figs. 2B-D). Although hypoxia increased autophagy within 24 hours, no significant changes in Prom1 were noted at this time point, indicating that sustained activation of autophagy in response to prolonged moderate hypoxia correlates with increased expression of Prom1. Furthermore, these results also suggest that Prom1 may play a direct or indirect role in the activation of autophagy.

Both mTORC1 and mTORC2 are negative regulators of autophagy, and phosphorylation of ribosomal protein S6 at Ser 235/236 (p-S6 Rp) and phosphorylation of Akt at Ser473 (p-Akt) are surrogate markers of mTORC1 and mTORC2 activation, respectively.26,47,48 Thus, we examined the activation of p-S6 Rp and Akt in RPE cells exposed to hypoxia. Hypoxia significantly decreased phosphorylation of p-S6 Rp at 48 hours, which was completely abolished at later time points. Furthermore, hypoxia decreased phosphorylation of p-Akt after 48 to 96 hours of hypoxia (Figs. 2B, 2D). These results suggest, markers of mTORC1 and mTORC2 activities are inhibited by hypoxia.

Amino acid starvation is a potent inducer of autophagy. Amino-acid deprivation of ARPE-19 cells by EBSS rapidly increased levels of LC3-II, Atg5, and Atg7 within 3 hours (Figs. 3A-C). After 6 to 9 hours of EBSS treatment, LC3-I levels decreased but LC3-II levels were higher compared to untreated cells. Both LC3-I and LC3-II levels were significantly reduced after 18 hours of EBSS treatment, due to lack of amino acid in the culture medium and sustained induction of autophagy. Similar to the effects of hypoxia on LC3 levels, LC3-II expression did not progressively increase in ARPE-19 cells treated with EBSS for 6 to 18 hours. Thus, normalization of LC3-II with actin in response to EBSS treatment cannot be used to demonstrate activation of autophagy in ARPE-19 cells. Because LC3-I levels decreased with time in ARPE-19 cells (Fig. 3A) and intense autophagy flux has been associated with consumption of LC3-II protein,49 we used LC3-II to LC3-I ratio to monitor changes in autophagy. The ratio of LC3-II to LC3-I showed time-dependent induction of autophagy in nutrient-deprived ARPE-19 cells (Fig. 3B). EBSS treatment increased phosphorylation of p62 Ser349 and its expression at 3 to 6 hours and gradually declined thereafter, indicating sustained activation of autophagy after prolonged amino-acid deprivation (Figs. 3A-C). Similar to the effects of hypoxia, EBSS inhibited Akt activation after 6 hours and decreased p-S6 Rp within 3 hours (Figs. 3A-C), which was completely blocked thereafter, indicating complete inhibition of mTORC1 and mTORC2 activities in response to nutrient deprivation. Importantly, Prom1 expression significantly increased between 3 and 6 hours of nutrient deprivation, which later returned to levels seen in control cells (Figs. 3A, 3C). To confirm the correlation of Prom1 with autophagic activity, we used flow cytometry, which was used successfully for assessing autophagy in cultured cells.50 ARPE-19 cells treated with EBSS for 3 hours showed significant increase of endogenous Prom1 (CD133) intensity, which correlated with increased intensity of LC3 (Figs. 3D, 3E) by flow cytometry. These data demonstrate that transient upregulation of Prom1 is associated with autophagy execution in response to amino acid deprivation.

Autophagy proteins and autophagy flux show an age-related increase in human and mouse RPE.51 Since metabolic stress (by nutrient deprivation) and hypoxia increase Prom1 expression and simultaneously induce autophagy in the RPE, we investigated whether age-related induction of autophagy was also correlated with Prom1 expression. RPE cells from aged donors show increased LC3-II/LC3-I ratio, Atg5 expression, reduced p62 expression, and increased formation LC3-II puncta (Figs. 4A-D), demonstrating age-related induction of autophagic activity relative to young RPE. Importantly, aged RPE cells also contain higher levels of Prom1 compared to young RPE. To further investigate whether aging activates autophagy flux, young and aged RPE cells were treated with BAF and CQ. CQ is a lysosomotropic agent that was used to inhibit lysosomal enzymes, which in turn leads to the inhibition of both fusion of autophagosomes with lysosomes and lysosomal degradation.51 Both BAF and CQ increased LC3-II levels in young RPE cells (Fig. 4E). In aged RPE cells, LC3-II/actin ratio significantly increased in response to BAF and CQ treatment when compared to young RPE cells (Figs. 4E, 4F) demonstrating that aging increases autophagy flux. Together, our results suggest a strong relationship between aging, Prom1 expression, and the induction of autophagy in the RPE.
Overexpression of WT Prom1 Induces Autophagy via Inhibition of mTORC1 and mTORC2 Activities

Having shown that increased Prom1 expression is correlated with the induction of autophagy, we investigated whether overexpression of Prom1 was able to induce autophagy in the RPE. Since primary RPE cells senesce with cell passage, we used the well-characterized human ARPE-19 cell lines for overexpression of Prom1. We infected ARPE-19 cells with a lentivirus that overexpresses Prom1 (WT) and selected for stably infected cells. WT cells have a robust increase in Prom1 protein (Fig. 5A) and mRNA expression (Fig. 5D), and increased expression of Atg5, Atg7, and LC3-I/LC3-II in confluent ARPE-19 cells exposed to EBSS for the indicated time periods. (B) LC3-I/LC3-II ratio and p62 densitometric analysis of results shown in A. *Significantly different compared to 0-hour cells (P < 0.05). (C) Densitometric analysis of results shown in A. Zero-hour values were set at 100%. *Significantly different compared to 0-hour cells (P < 0.05). (D) Flow cytometry analysis demonstrating CD133 intensity (antibody from Miltenyi) in response to EBSS treatment for 3 hours in ARPE-19 cells. (E) Flow cytometry analysis showing LC3 intensity after 3 hours of EBSS treatment in ARPE-19 cells.

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previously for assessing autophagy flux in cultured cells.\textsuperscript{42} Optimization of image analysis by the IDEAS software and the spot counting wizard (described in the Methods section) allowed us to evaluate Prom1 expression and LC3 spot count in control cells and cells overexpressing Prom1 (Fig. 5E). Our analyses showed that overexpression of Prom1 increased intensity of CD133 (Prom1) green staining (Fig. 5E). Cells overexpressing Prom1 significantly increased basal LC3 spot count $>3$ in single cells (red staining, Figs. 5E, 5F). To demonstrate changes in autophagic flux, WT cells were treated in the presence and absence of CQ for 3 hours. CQ increased LC3 spot count $>3$ in control ARPE-19 cells, which further increased significantly in WT cells (Fig. 5G), demonstrating increased autophagic flux.

Although several Prom1 isoforms are expressed in the retina,\textsuperscript{52} it is unclear which splice variant is expressed in the RPE. Using RT-PCR, we analyzed expression of Prom1 mRNA transcript variant 3 in the RPE. Since Prom1 retina isoforms differ on the presence or absence of exons,\textsuperscript{52} specific primers for the amplification of various exons of transcript variant 3 were used. In control ARPE-19 cells, mRNA was detected from exons-2, 3, 4, 6, and exons 27 and 29 from the UTR region.
Since the lentiviral Prom1 construct used to generate the WT cells contained the cDNA sequence ranging from 236 to 2806 bp, mRNAs from all exons of Prom1 transcript variant 3 were detected in these cells except for exons 27 and 29 from the UTR region (PCR-4). This absence is a negative control, showing the UTR region (exon 27 and 29) is after the stop codon in WT cells. Sequence analysis of the purified PCR products (data not shown) confirmed that lentivirus-mediated overexpression of Prom1 (WT) results in the expression of the same transcript variant 3.

In order to confirm the effects of Prom1 on autophagic flux and LC3 processing, both control cells and WT cells were treated with BAF for various time periods. The ratio of LC3-II to LC3-I is regarded as a critical indicator of autophagic activation in RPE cells.\(^{27}\) In control cells, 100 nM BAF increased LC3-II/LC3-I ratio from 1 to 3 hours, and robustly increased p62 expression from 1 to 3 hours (Figs. 6A, 6B). In WT cells, BAF...
further increased LC3-II/LC3-I ratio in a time-dependent manner (Figs. 6A, 6B). However, the levels of p62 accumulation in response to BAF were significantly lower compared to control cells, suggesting increased autophagic flux and rapid LC3-II consumption in WT cells.

We next verified if autophagic flux was potentiated in response to Prom1 overexpression. Control ARPE-19 and WT cells were cultured in the presence or absence of BAF in normal growth media (FBS) or amino acid starvation media (EBSS). Consistent with data presented in Figure 6A, BAF increased LC3-II accumulation in control cells. EBSS alone decreased LC3-I and partially increased LC3-II, whereas EBSS+BAF decreased LC3-I but robustly increased LC3-II expression in control cells. In Prom1 overexpressing WT cells, BAF alone increased LC3-II, which was higher compared to control ARPE-19 cells. Interestingly, EBSS decreased LC3-I and increased LC3-II levels in WT cells, and BAF further increased LC3-II levels in EBSS treated WT cells. Because LC3-II levels rapidly increased in WT cells, we used LC3-II/actin ratio and p62/actin ratio to measure autophagy in WT cells. WT cells under normal growth conditions have increased LC3-II/actin ratio, and decreased levels of p62/actin ratio, phosphorylation of Akt, and S6 Rp showing constitutive and sustained activation of autophagy. This response was enhanced by BAF treatment (Figs. 6C, 6D). EBSS treatment in WT cells further increased LC3-II/actin ratio and decreased p62, p-Akt, and p-S6 Rp expression, confirming potentiation of autophagy. An enhanced response was observed for WT cells treated with
EBSS+BAF (Figs. 6C, 6D), demonstrating that Prom1 overexpression potentiates autophagy flux in response to BAF and amino acid starvation.

Next, we compared the correlation between mTOR activity and the induction of autophagy in control (parent ARPE-19) and WT cells using three different mTOR inhibitors, RAP, or both Torin1 and Torin 2. RAP is an allosteric inhibitor of mTOR and partly suppresses mTORC1 function, whereas both Torin 1 and Torin 2 are catalytic inhibitors and are capable of completely blocking both mTORC1 and mTORC2 activities.53,54 Consistent with our previous observations (Fig. 6), BAF increased both p62 and LC3-II/actin ratio mainly due to the impaired fusion of autophagosomes with lysosomes. Compared to control cells at the corresponding treatment, WT cells have decreased p62, and p-S6 Ribosomal protein Ser235/236 expression, indicating activation of autophagy (Figs. 7A–C). This trend was observed for all treatment groups and was further enhanced when cells were co-treated with BAF. Notably, BAF significantly increased LC3-II/actin ratio in WT cells co-treated with RAP and Torins, demonstrating increased autophagic flux.

**Figure 7.** mTOR inhibitors potentiate autophagy flux in cells overexpressing Prom1 and enhancement of autophagy confers protection from Nutlin-3-induced apoptosis. (A) Representative immunoblots showing the levels of LC3-I/LC3-II, p-Akt Ser473, total-Akt, p-S6 Ribosomal protein Ser235/236, and p62 in control and Prom1 overexpressing ARPE-19 cells (WT) treated with 1.5 μM RAP or Torins (T: 1.5 μM Torin1 plus 3 μM Torin2) in the presence and absence of 100 nM BAF for 3 hours. (B) Densitometric analysis of LC3-II and p62 data presented in A. *Significantly different compared to untreated (UT) control cells (P < 0.05); #Significantly different compared to corresponding control group (P < 0.05). (C) Densitometric analysis of results presented in A. Values from control untreated (UT) cells were set at 100%. *Significantly different compared to UT cells (P < 0.05); #Significantly different compared to corresponding control group (P < 0.05). Overexpression of Prom1 induces autophagy but inhibits apoptosis. (D) Data showing LC3-II/actin ratio, active caspase-3, and Atg5 levels in control and Prom1 overexpressing cells treated with varying doses of Nutlin-3 for 3 hours. (E) Densitometric analysis of results presented in A. *Significantly different compared to cells untreated with Nutlin-3 (P < 0.05). #Significantly different compared to control cells treated with or without Nutlin-3 (P < 0.05).
Since the mTOR inhibitors+BAF potentiated LC3-II/actin ratio in WT cells, we measured Akt and p-S6 phosphorylation. Basal levels of Akt/total Akt ratio were low in WT cells and treatment of WT cells with RAP, Torins, or Torins+BAF failed to significantly decrease p-Akt/total Akt ratio compared to control cells. These mTOR inhibitors significantly reduced low basal levels of p-S6 Rp phosphorylation in WT cells in the presence or absence of BAF. Overall, expression of p62 in WT cells treated with mTOR inhibitors in the presence and absence of BAF were considerably lower compared to control cells, indicating increased autophagic flux (Figs. 7A, 7B). Interestingly, BAF treatment in control cells decreased p-Akt/total Akt ratio and increased p-S6 Rp/actin ratio, indicating different roles of BAF on mTORC1 and mTORC2 signaling. This difference was observed for p-S6 Rp to a lesser extent in WT cells treated with BAF alone. Together, these results demonstrate that Prom1 regulates RPE autophagy by negatively regulating both mTORC1 and mTORC2 activities.

To further confirm whether Prom1-dependent RPE autophagy confers cytoprotection, we treated control cells and WT cells with different doses of the nongenotoxic p53 activator, Nutlin-3. Consistent with our previous studies, Nutlin-3 had no effect on ARPE-19 apoptosis, but 60 μM Nutlin-3 sensitized cells to apoptosis as evidenced by high levels of caspase-3 activation (Fig. 7D). Concomitant with caspase-3 activation, both 40 μM and 60 μM Nutlin-3 significantly increased LC3-II/LC3-I ratio (Fig. 7E) suggesting that Nutlin-3 is a potent inducer of apoptosis, which in turn induces autophagy in control cells. Nutlin-3 further potentiated autophagy in WT cells in a dose-dependent manner as seen by the higher levels of Atg5 expression and the increase in LC3-II/actin ratio. Importantly, Nutlin-3 failed to activate caspase-3 in cells overexpressing Prom1, indicating that increased autophagic flux confers protection from Nutlin-5-induced apoptosis. The lack of caspase-3 activation in cells overexpressing Prom1 fails to induce proteolytic cleavage of LC3. As a result, the LC3-II/actin ratio in cells overexpressing Prom1 and treated with Nutlin-3 is higher compared to control cells. These results demonstrate that Prom1-dependent induction of autophagy in the RPE does not permit induction of apoptosis in response to nongenotoxic stress.

**CRISPR-Mediated Prom1 KO Activates mTOR Signaling and Impairs Autophagy Flux**

To further demonstrate the central role of Prom1 in regulation of autophagy, we used Prom1 KO ARPE-19 cell lines (hereafter referred to as KO or KO-6) using the CRISPR-Cas9 lentiviral construct as described in the Methods section. The use of innovative CRISPR technology allows genome editing of the genetic code, typically causing a KO or complete elimination of gene function. We used the 17-nucleotide guide-RNA (gRNA) sequence to target Prom1 gene at exon 10, and selected guides were cloned into the lentiviral backbone. Cas9 lentiviral construct as described in the Methods section. (heretofore referred to as KO or KO-6) using the CRISPR-Cas9 lentiviral construct as described in the Methods section. To further elucidate the involvement of the mTOR/Akt signaling axis in Prom1-mediated autophagy, we treated control Cas9 and KO cells with Torins (using Torin1 and Torin 2) in the presence or absence of BAF. Torins increased Prom1, LC3-II, and decreased p62 expression in control Cas9 cells due to inhibition of mTOR signaling and increased synthesis of autophagosomes. Control cells treated with Torins in the presence of BAF (Torins+BAF) showed higher levels of Prom1, suggesting that autophagy induction in response to mTOR inhibition (by Torins) correlates with increased expression of Prom1 (Figs. 9A, 9B). Higher LC3-II and p62 levels were observed in response to Torins+BAF, indicating that accumulation of LC3-II is due to inhibition of autophagosomal cargo degradation. Torins failed to increase Prom1, LC3-II in KO cells, and lower levels of LC3-II were observed in KO cells treated with Torins+BAF (Figs. 9A, 9B). Unlike WT cells, LC3-I levels remained unaltered in the KO cells after treatment with Torins and Torins+BAF. To compare the LC3 data obtained with WT cells (Figs. 5–7), we used LC3-II/LC3-I to evaluate autophagy in KO cells. Untreated KO cells have decreased LC3-II/LC3-I ratio, and Atg5 expression, and increased p62, p-Akt, and p-S6 Rb expression, suggesting KO cells have increased mTORC1/2 signaling and decreased basal autophagy (Figs. 9A, 9B). Treatment of KO cells with Torins showed significantly lower LC3-II/LC3-I ratio and Atg5, but failed to decrease p62, when compared to control cells. Combined Torins+BAF treatment in KO cells had decreased LC3-II/LC3-I and Atg5 expression, when compared to control cells. Consistent with our previous observations (Fig. 8A), Prom1 KO cells have high basal levels of p-Akt and p-S6 Rb, indicating basal activation of mTORC1 and mTORC2 signaling. However, Torins treatment in KO cells completely inhibited both p-Akt and p-S6 Rp levels (Figs. 9A, 9B) but had no effect on Atg5 expression, showing that inhibition of Akt and S6-Rp is insufficient to induce autophagic activity in the absence of Prom1. Interestingly, KO cells have high levels of p62, which was not altered by either Torins or Torins+BAF treatment, confirming that lack of Prom1 inhibits autophagic flux due to decreased autophagosome maturation and fusion of autophagosomes with lysosomes.

To demonstrate this, we examined LC3 puncta formation in Prom1 KO cells treated with 50 μM CQ (3 hours) or EBSS (3 hours) by confocal microscopy. Since amino-acid starvation by EBSS inhibited mTORC1 and mTORC2 activities leading to the induction of autophagy in ARPE-19 cells (Figs. 3A–C), we used EBSS in control and Prom1 KO cells. Using confocal microscopy, co-localization of LC3 puncta with a lysosomal marker, lysotracker, was used to detect the trafficking of autophagosomes to lysosomes. KO of Prom1 (clone-6) significantly decreased the number of LC3+ puncta per cell at baseline and after CQ or EBSS treatment (Figs. 9C, 9D). Co-localization of LC3 aggregates with lysotracker dramatically increased in control cells in response to CQ or EBSS treatment. However, the extent of puncta-lysosome co-localization in response to EBSS was lower compared to CQ-treatment in control cells, showing that CQ specifically inhibits lysosomal degradation, which in turn increases the accumulation of LC3 puncta in lysosomes. The EBSS-induced LC3 puncta was effectively cleared by lysosomal degradation, which prevented its accumulation in control cells. KO of Prom1 (clone-6) significantly decreased co-localization of LC3 puncta with lysosomes after EBSS or CQ treatment (Figs. 9C, 9D). To further validate our observations, we compared LC3 puncta formation...
in control and Prom1 KO cells treated with EBSS (4 hours) or 200 nM RAP (4 hours) by confocal microscopy. Both RAP and EBSS induce autophagy through mTOR inhibition in ARPE-19 cells. These activators of autophagy increased LC3 puncta per cell and their co-localization to lysosomes in control Cas9 cells, which was significantly blocked in KO cells (Supplementary Fig. S2), further confirming that Prom1 is required for biogenesis of autophagosomes and delivery of the autophagosomal cargo to lysosomes in the RPE.

To further elucidate whether stress-dependent induction of autophagy requires the participation of Prom1, we treated control and KO cells with EBSS medium for the indicated time periods (Fig. 10A). Consistent with our previous observations (Fig. 3A), amino-acid deprivation by EBSS significantly increased Prom1 expression at 3 hours in Cas9-infected control cells. Furthermore, EBSS rapidly induced autophagy within 3 hours and sustained activation of autophagy was evident by time-dependent decrease of LC3-I and increase of LC3-II/LC3-I ratio in control cells (Figs. 10A, 10B). Both LC3-I and LC3-II levels were dramatically reduced by 18 hours indicating consumption of LC3 proteins due to sustained activation of autophagy. KO of Prom1 impaired EBSS-induced autophagy,
which was evident by unaltered levels of both LC3-I and LC3-II at 3 hours, lower LC3-II levels at 18 hours, and lower LC3-II/LC3-I ratio from 0 to 18 hours compared with control-Cas9 cells. Prom1 KO decreased basal expression of Atg5 and Atg7, and EBSS failed to alter expression of these autophagy-related proteins suggesting that loss of Prom1 decreases autophagic activity. KO cells at 0 hours had high basal levels of p-Akt and p-S6 Rb, which were inhibited or abolished, respectively, in response to EBSS treatment. Although EBSS inhibited these key markers of mTOR signaling in KO cells, it failed to restore autophagy. This indicates that Prom1 deletion constitutively activates mTORC1 and mTORC2 and that inhibition of mTORC1 and mTORC2 activities is insufficient to trigger autophagy in the absence of Prom1. Furthermore, p62 levels remained elevated throughout the entire time period of EBSS treatment strongly (Figs. 10A, 10B) reinforcing the observation that absence of Prom1 impairs normal trafficking of autophagosomes with lysosomes.

Multispectral flow cytometry was also used to demonstrate a role of Prom1 in the biogenesis of autophagosomes in response to amino-acid starvation. Control and KO cells were treated with EBSS for 3 hours and stained with Prom1/CD133.
and LC3-I/II specific antibodies. Double positive cells, single positive CD133$^+$ cells, LC3-I/II$^+$ cells, and LC3-I/LC3-II in Cas9 and Prom1-KO ARPE-19 cells treated with EBSS for the indicated time periods. (B) Densitometric analysis of data presented in A. *Significantly different compared to control cells at 0 hours ($P < 0.05$). #Significantly different compared to respective control cells ($P < 0.05$). (C) Representative BF, CD133$^+$, LC3$^+$ images by FlowSight Imaging Flow Cytometer in control and Prom1-KO cells in the presence and absence of EBSS for 3 hours. (D) % Total LC3 spot count of 3 or greater in control and KO cells in the presence of EBSS relative to untreated (UT) cells. *Significantly different compared to control cells treated with EBSS ($P < 0.05$).

**Figure 10.** KO of Prom1 impairs starvation-induced autophagy. (A) Representative immunoblots showing the levels of Prom1, p-Akt Ser473, Atg5, Atg7, p-S6 Rbp, Ser235/236, p62, and LC3-I/LC3-II in Cas9 and Prom1-KO ARPE-19 cells treated with EBSS for the indicated time periods. (B) Densitometric analysis of data presented in A. *Significantly different compared to control cells at 0 hours ($P < 0.05$). #Significantly different compared to respective control cells ($P < 0.05$). (C) Representative BF, CD133$^+$, LC3$^+$ images by FlowSight Imaging Flow Cytometer in control and Prom1-KO cells in the presence and absence of EBSS for 3 hours. (D) % Total LC3 spot count of 3 or greater in control and KO cells in the presence of EBSS relative to untreated (UT) cells. *Significantly different compared to control cells treated with EBSS ($P < 0.05$).

**Prom1 Associates With p62 and Is Required for Autophagosome-Lysosome Fusion**

Considering our finding that KO of Prom1 constitutively activates mTORC1 and mTORC2, and inhibition of mTORC1/2 fails to restore autophagic flux in these cells, we hypothesized a significant association between Prom1 and other protein components of the autophagy machinery. To examine this possibility, we immunoprecipitated Prom1 from control ARPE-19 cells and WT cells. p62 was detected in the Prom1 immune complexes from control (both C and empty vector EV infected) cells and at significantly higher levels in WT cells, suggesting that Prom1 interacts with p62, and ectopic overexpression of Prom1 enhances the presence of p62 in the Prom1 immunoprecipitates (Figs. 11A, 11D). Reciprocal coinmunoprecipitation assays showed that p62 was bound to Prom1 in control cells, and WT cells have lower levels of Prom1 bound to p62 (Figs. 11B, 11D), mainly due to decreased p62 levels in WT cells. Prom1, p62, and HDAC6
bands were undetected in the mock immunoprecipitated samples. Input confirmed increased Prom1 and decreased p62 expression in WT cell extracts (Fig. 11C).

The ubiquitin-binding cytosolic deacetylase-6 (HDAC6) controls autophagy by regulating the fusion of autophagosome with lysosomes. Recent studies show that Prom1 interacts with cytosolic HDAC6, suggesting the possibility that a macromolecular complex comprising of p62, HDAC6, and Prom1 may play a central role in autophagosome maturation and its fusion to the lysosomes. To test the formation of this macromolecular complex, we analyzed Prom1 immunoprecipitates for HDAC6. Endogenous Prom1 was found to associate with HDAC6, and overexpression of Prom1 (WT) increased the amount of HDAC6 in the Prom1 immunoprecipitates (Figs. 11A, 11D). Reciprocal immunoprecipitation with p62 showed the presence of HDAC6 in control cells, but overexpression of Prom1

**FIGURE 11.** Coimmunoprecipitation of Prom1, p62, and HDAC6. (A) Prom1 immunoprecipitates from untreated ARPE-19 control (UT), Cas9 empty lentivirus (EV), and Prom1 overexpressing (WT) cell lysates were analyzed for the presence of Prom1, p62, and HDAC6. The mixture of beads with the respective antibodies was used for mock immunoprecipitation. (B) p62 immunoprecipitates were analyzed for the levels of Prom1, HDAC6, and p62. (C) Equal amounts of whole cell lysates were simultaneously analyzed for the levels of Prom1, p62, and actin as input. (D) Quantification of proteins in Prom1 and p62 immunoprecipitates by densitometry. Control values were set at 100%. *Significantly different compared to untreated control cells (P < 0.05). (E) Prom1 immunoprecipitates from UT, EV, and Prom1-KO, and KO-6 cell lines were analyzed for the presence of Prom1, HDAC6, and p62. (F) Data showing the levels of Prom1, HDAC6, and p62 in p62 immunoprecipitates from control, EV, and KO samples. (G) Equal amounts of UT, EV, KO, and KO-6 cell lysates were simultaneously analyzed for the levels of Prom1, p62, and actin as input. (H) Densitometric analysis of immunoprecipitation data presented in D and E. *Significantly different compared to control cells (P < 0.05).
did not significantly increase the levels of HDAC6 in the p62 immunoprecipitates (Figs. 11B, 11D), showing that Prom1 is not required for p62-HDAC6 association.

In order to further confirm that Prom1 interacts with p62 and HDAC6, we probed control-Cas9 and Prom1-KO cell lines. Immunoprecipitation of Prom1 from control (C and EV) cells contained both Prom1 and p62. Prom1 was not visible in Prom1 immunoprecipitates from KO and KO-6 cells, and the levels of p62 were significantly lower in these immune complexes (Figs. 11E, 11H), showing that absence of Prom1 alters the association of p62 with Prom1. Consistent with previous observations (Figs. 8–10), KO of Prom1 significantly increased p62 expression (input, Fig. 11G). However, reciprocal p62 immunoprecipitates from KO cells contained undetectable levels of Prom1, but higher levels of p62 (Fig. 11F). Prom1 and p62 immunoprecipitates from cells both overexpressing and lacking Prom1 contained comparable levels of HDAC6, indicating that the status of cellular Prom1 does not impact the association of p62 with HDAC6 (Figs. 11D, 11H).

Our overall findings suggest that Prom1 is an important component of a macromolecular complex consisting of p62 and HDAC6. Disruption of the Prom1 association with this complex negatively impacts autophagosome maturation and delivery of autophagosomes to lysosomes.

**DISCUSSION**

The key finding of these studies is that Prom1 is a pivotal regulator of autophagy in the RPE. Our data demonstrate for the first time that overexpression of Prom1 constitutively activates autophagy in the RPE via inhibition of mTORC1 and mTORC2. Conversely, KO of Prom1 impairs RPE autophagy via upregulation of mTORC1/2 activities, suggesting that Prom1 is central to the regulation of autophagy. Our data suggest that this effect is controlled in part due to Prom1’s ability to form a macromolecular complex comprising of autophagy proteins p62 and cytosolic HDAC6, and disruption of this interaction by genetic deletion of Prom1 impairs autophagosome biogenesis and causes defects in trafficking of the autophagosomes to the lysosomes. These findings have important implications for the maintenance of RPE homeostasis because defective autophagosomal-lysosomal-phagocytic pathways can lead to ineffective clearance of shed photoreceptor outer segments and cause accumulation of damaged organelles and protein aggregates including lipofuscin-like debris in lysosomes, all of which have been linked with the pathogenesis of age-related retinal diseases, including AMD.

The connection between autophagy and aging is complex and cell-type specific. For instance, aging alone has been implicated in both promoting and inhibiting autophagy in various cell types. Age-related deterioration of vision has been linked to the decline in noncanonical autophagy coupled with a loss of phagocytic activity in the RPE, suggesting that the functional interplay between autophagy and phagocytosis is fundamental to vision. In keeping with this notion, our data show that aging increased classical autophagy in the RPE, supported by increased expression of Atg5, loss of p62 expression, increased levels of autophagic puncta, and increased autophagic flux. Of note, age-related induction of autophagy in the RPE was associated with increased expression of Prom1. Therefore, the induction of Prom1-dependent autophagy during aging may be an intrinsic defense mechanism, which enables the RPE to cope with increased oxidative burden, accumulation of ubiquitinated/nonubiquitinated protein aggregates, and phagocytic activity. Although Prom1 was initially described as a surface antigen in normal hematopoietic stem cells and cancer stem cells, recent studies have demonstrated the presence of Prom1 in normal adult tissues including the retina suggesting that Prom1 has diverse physiological functions beyond the known association with cancer stem cells. Several Prom1 isoforms are expressed in the retina, but there is no Prominin-2 (Prom-2) expression in the RPE, which perhaps explains why a Prom1 mutation in the human gene causes retinal degeneration without causing other pathological abnormalities. Prom1’s important structural role in the photoreceptor outer segment membrane suggested a similar structural role of the protein in the RPE apical microvilli. Anti-CD133 antibody staining of tissue sections showed expression patterns of Prom1 in the apical surface of the RPE layer, the interface between the neural and epithelial retina, and at the basal part of the outer segment of rods and cones in the adult mouse. Surprisingly, our studies with human cells in vitro demonstrated that Prom1 is expressed in the RPE, but the protein is primarily cytosolic. Thus, the primary role of Prom1 in the human RPE is not structural, and there are differences in spatiotemporal expression and localization of Prom1 between in vitro cultures and in vivo tissue sections. Since Prom1 mutations cause dominant macular degeneration in humans and mice, it is likely that Prom1 is an essential component of the intracellular membrane-enclosed organelles, including ER and Golgi bodies. Several models have attempted to explain the subcellular localization of proteins, including mono and poly-ubiquitination of proteins. Since ubiquitination of Prom1 has been reported, it is attractive to speculate that ubiquitination regulates cytoplasmic localization of Prom1 in the RPE. A recent study showed that Prom1 was dynamically released from plasma membrane into cytoplasm in response to high glucose, raising the possibility of Prom1’s cytoplasmic accumulation by its trafficking from the cell surface. However, only few studies have reported the intracellular localization of Prom1 in addition to its cell surface expression.

To elucidate the role of Prom1 in the RPE, we performed a series of studies using overexpression and genomic editing strategies. These studies were aimed at spatiotemporal modulation of Prom1 expression, which could impact RPE function. Lentiviral overexpression of Prom1 potentiates expression of autophagy markers Atg5, Atg7, decreased p62 accumulation, and constitutively activated autophagy. Furthermore, overexpression of Prom1 resulted in the expression of the same splice variant, which is endogenously expressed in ARPE-19 cells, demonstrating that our observations are focused on one splice variant of Prom1 and its function. To rule out the involvement of other possible Prom1 splice variants in the regulation of autophagy in ARPE-19 cells, we used CRISPR/Cas9-mediated genome editing, which is capable of deleting all splice variants. Genomic deletion of Prom1 blocked basal Atg5, Atg7 expression, increased p62 expression, and, consequently inhibited basal autophagy flux, reinforcing the concept that Prom1 is a central regulator of autophagy in vitro and in vivo. It is unclear how Prom1 regulates the expression of Atg5 and Atg7 in the RPE and warrants further investigation. The RPE layer is well preserved in 1-month-old Prom1−/− mouse
Prom1 Negatively Regulates mTOR Signaling

Prom1, a receptor tyrosine kinase, has been shown to negatively regulate mTOR signaling. In agreement with this, a recent study demonstrated that Prom1-mediated upstream suppression of mTORC1/2 is mechanistically linked to downstream events in the autophagy pathway. Increased Prom1 expression led to upstream inhibition of both mTORC1 and mTORC2. These cells also exhibited reduced LC3 puncta formation and decreased trafficking of autophagosomes with lysosomes, suggesting that Prom1 plays a role in mediating upstream and downstream roles in autophagy and cellular homeostasis in the RPE.

Hypoxia, nutrient deprivation, and ER stress upregulate autophagy, an adaptive housekeeping mechanism that promotes organismal health and counteracts the aging process. Cells exposed to stressors utilize various strategies to survive of diverse cell types, including cancer stem cells. Further, Prom1 trafficking in response to changes in cell microenvironment such as low glucose and hypoxia caused release of membrane bound Prom1 to the cytoplasm, which promotes autophagy in cancer cells, suggesting that dynamic alteration of CD133 localization from membranes to the cytosol in response to diverse environmental cues is necessary for the induction of autophagy. In hepatoma cells, expression of Prom1/CD133 promoted autophagosome formation, and silencing of Prom1 attenuated this activity further confirming that Prom1 is an essential component of the autophagic machinery.

In agreement with this conclusion, we propose a conceptual model in which increased Prom1 expression and genetic deletion of Prom1 abrogated autophagosome maturation and basal autophagy in primary RPE cultures. We examined whether exposure of RPE cells to various stressors including hypoxia and nutrient deprivation triggered autophagic activity with concomitant upregulation of Prom1 expression. Relative hypoxia increased autophagy protein Atg5 and decreased p62 expression with concomitant reduction of its phosphorylation at Ser349, indicating robust induction of autophagy flux. It is noteworthy that induction of autophagy was coupled with increased Prom1 expression after prolonged exposure to hypoxia, suggesting that Prom1 protects the RPE against prolonged hypoxia by upregulation of autophagy. Exposure of RPE cells to EBSS medium rapidly activated autophagy. More importantly, EBSS-mediated induction of autophagy is correlated with increased expression of Prom1, suggesting that nutrient deprivation-induced autophagosome formation requires the engagement of Prom1. Mechanistically, mTOR is a central signaling pathway that coordinates the cellular processes with metabolic homeostasis through its ability to negatively regulate autophagy. Both nutrient deprivation and hypoxia rapidly inactivated mTORC1. These stress signals had a similar effect on mTORC2 inhibition. Together, these results suggest a model in which increased Prom1 expression and mTORC1/mTORC2 inhibition orchestrate the induction of autophagy flux in response to stress signals in the RPE (Fig. 12).

We analyzed the mechanisms through which Prom1 regulates autophagy. Although mTORC1 and mTORC2 are independent regulators of autophagy, mTORC2 can both positively and negatively regulate autophagy. The importance of this process was verified in Prom1 KO cells that showed increased activation of mTORC1 and mTORC2. These cells also exhibited reduced LC3 puncta formation and decreased trafficking of autophagosomes with lysosomes, suggesting that Prom1 plays an important upstream and downstream role in autophagy and cellular homeostasis in the RPE.

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molecular events increased p62 accumulation, impaired autophagosome maturation, and autophagy flux. Together, our results demonstrate that Prom1 is a key regulator of autophagy in the RPE. Modulation of its age-related expression may represent a cellular mechanism that protects the cell from age-related decline in autophagy flux, a critical role played by the RPE. It is interesting to speculate that such a decline in cellular homeostasis may play a mechanistic role in the AMD phenotype and Stargardt-like (STGD4) macular dystrophy disease. In addition, the newly defined role for Prom1 in autophagy may explain the resistance of CD133+ cancer cells to hypoxia and chemotherapy.

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