Prevention of Proliferative Vitreoretinopathy by Suppression of Phosphatidylinositol 5-Phosphate 4-Kinases

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Submitted: February 23, 2016
Accepted: June 9, 2016

PURPOSE. Previous studies have shown that vitreous stimulates degradation of the tumor suppressor protein p53 and that knockdown of phosphatidylinositol 5-phosphate 4-kinases (PI5P4K and β) abrogates proliferation of p53-deficient cells. The purpose of this study was to determine whether vitreous stimulated expression of PI5P4K and β and whether suppression of PI5P4K and β would inhibit vitreous-induced cellular responses and experimental proliferative vitreoretinopathy (PVR).

METHODS. PI5P4K and β encoded by PIP4K2A and 2B, respectively, in human ARPE-19 cells were knocked down by stably expressing short hairpin (sh)RNA directed at human PIP4K2A and -2B. In addition, we rescued expression of PI5P4K and β by re-expressing mouse PIP4K2A and -2B in the PI5P4K and β knocked-down ARPE-19 cells. Expression of PI5P4K and β was determined by Western blot and immunofluorescence. The following cellular responses were monitored: cell proliferation, survival, migration, and contraction. Moreover, the cell potential of inducing PVR was examined in a rabbit model of PVR effected by intravitreal cell injection.

RESULTS. We found that vitreous enhanced expression of PI5P4K and β in RPE cells and that knocking down PI5P4K and β abolished vitreous-stimulated cell proliferation, survival, migration, and contraction. Re-expression of mouse PIP4K and -β in the human PI5P4K and β knocked-down cells recovered the loss of vitreous-induced cell contraction. Importantly, suppression of PI5P4K and β abrogated the pathogenesis of PVR induced by intravitreal cell injection in rabbits. Moreover, we revealed that expression of PI5P4K and β was abundant in epiretinal membranes from PVR grade C patients.

CONCLUSIONS. The findings from this study indicate that PI5P4K and β could be novel therapeutic targets for the treatment of PVR.

Keywords: proliferative vitreoretinopathy, PI5P4K, vitreous

Proliferative vitreoretinopathy (PVR) develops in 8% to 10% of patients who undergo reparative primary retinal detachment surgery1-7 and in 40% to 60% of patients with open globe injury.18-20 The major feature of PVR is the formation of a subretinal or epiretinal membrane (ERM) that consists of retinal pigment epithelial (RPE) cells, fibroblasts, glial cells, and macrophages, as well as extracellular matrix.2-7 The ERM may be attached to the retina and subsequently contract, causing a new retinal detachment or failure of a surgically corrected detachment.2-7 At present, repeat surgery is the only option for treating PVR; however, the surgery has had poor functional results.

We have recently shown that vitreous from rabbits and patients with PVR preferentially activates the signaling pathway of PDGF receptor (PDGFRα)/phosphoinositide 3 kinase (PI3K)/Akt,17 leading to the phosphorylation of murine double minute 2 (MDM2), an oncogene protein acting as an E3 ubiquitin ligase.18-20 Phosphorylation of MDM2 increases its interaction with the tumor suppressor protein p53, promoting p53 degradation.21 The activation of p53 can lead to cell cycle arrest, apoptosis, and/or senescence.22 The codon 72 polymorphisms (rs1042522) in human p53 protein contain two variants, Arg and Pro. The Arg 72 variant is associated with more apoptosis than is the Pro 72 variant.23 The Arg 72 variant also interacts more readily with MDM2,24,25 which facilitates the export of p53 from the nucleus and its degradation through the ubiquitination pathway. Intriguingly, the Pro variant of p53 codon 72 polymorphisms is associated with a higher risk of developing PVR after a primary retinal detachment.25 Nutlin-3, a small molecule that blocks the
interaction of MDM2 with p53, prevents retinal detachment in experimental PVR in rabbits induced by intravitreal cell injection, but the cellular membranes still form in the vitreous. In addition, vitreous-induced cell contraction is stronger than that of cells with p53 suppressed by short hairpin (sh) RNA and without rabbit vitreous (RV) induction. Thus, it appears that other factors coordinate with p53 loss to promote vitreous-stimulated gel contraction as well as membrane formation in experimental PVR. Knowing that suppression of phosphatidylinositol 5-phosphate 4-kinases (PI5P4Ks) inhibits p53-null tumor growth, we hypothesized that PI5P4Ks might participate in vitreous-induced cellular responses as well as in the pathogenesis of PVR. Phosphatidylinositols (PIs), a family of phosphatidylglycerides, are synthesized by the phosphorylation at positions 3, 4, and 5 on the inositol ring. PI-4,5-bisphosphate (PI(4,5-P2) is the major substrate for class I PI3Ks and plays a key role in mediating the localization of proteins to the plasma membrane. The type II PI-5-P-4 kinases can catalyze the phosphorylation of PI(4,5-P2) at the 4-position to become PI-4,5-P2. In humans and mice, there are three distinct type II kinases, PI5P4Kα, β, and γ, encoded by PIPK2A, B and C, respectively. The type I PIP kinases generate PI4,5-P2 at the plasma membrane, whereas the type II kinases are located at internal membranes, including the endoplasmic reticulum, Golgi, and nucleus, where they likely generate PI(4,5-P2).

The localized type II PI5P4Ks may be involved in the development of PVR, and in contraction as well as the development of experimental PVR in rabbits. In addition, both PI5P4Kα and -β are abundant in the ERMs of grade C PVR patients.

Materials and Methods

Major Reagents and Cell Culture

Primary antibodies against p53, PI5P4Kα, and PI5P4Kβ were purchased from Cell Signaling Technology (Danvers, MA, USA) and -β-actin from Santa Cruz Biotechnology (Santa Cruz, CA, USA). Secondary antibodies, HRP (horseradish peroxidase)-conjugated goat anti-rabbit IgG and goat anti-mouse IgG, were purchased from Santa Cruz Biotechnology. Enhanced chemiluminescent substrate for detection of HRP was obtained from Pierce Protein Research Products (Rockford, IL, USA). Puromycin and carbencicillin were purchased from Sigma-Aldrich Corp. (St. Louis, MO, USA).

ARPE-19 (a spontaneously arising RPE cell line) cells were purchased from American Type Culture Collection (ATCC; Manassas, VA, USA), and RPEM cells were RPE cells originated from human RPE (ARPE-19) cells inhibited by Western blot.32–34 Cells Whose PI5P4Kα and β Were Knocked Down by shRNA

Escherichia coli strains containing plasmids pLKO.1-shRNA for PIPK2A (TRCN000006009 and TRCN000006011) and PIPK2B (TRCN000006013 and TRCN000006016) were purchased from Dharmaco (Lafayette, CO, USA), and an E. coli strain containing pLKO.1-shRNA for green fluorescent protein (GFP) was obtained from Dana-Farber Cancer Institute (Boston, MA, USA). The respective plLKO.1-shRNA lentiviral vector (1000 ng), the packaging plasmid psPAX2 (12260; Addgene, Cambridge, MA, USA) (900 ng), and the envelope plasmid pVS-V (Addgene 8454) (100 ng) were mixed together and then added to a mixture of TransIT-LT1 (MIR 2300; Mirus Bio LLC, Madison, WI, USA) or lipofectamine 2000 (Invitrogen) 6 µL with OPTI-MEM (Invitrogen 31985-070) 90 µL. This transfection mix was incubated at room temperature for 30 minutes and then carefully transferred into a 60-mm cell culture dish with HEK 293T cells that were approximately 70% confluent without antibiotics. After 18 hours (37°C, 5% CO2), the medium was replaced with growth medium supplemented with 30% FBS, and lentiviruses were harvested at 40 hours after the transfection. The viral harvest was repeated at 24-hour intervals three times. The virus-containing media were pooled and then centrifuged at 800g for 5 minutes, and the supernatant was used to infect cells of ARPE-19, RPEM, and RCFs supplemented with 8 µg/mL polybrene. The infected cells were selected in media with puromycin (ARPE-19, RPEM: 4 µg/mL; RCF: 2 µL/mL), and the resulting cells were examined by Western blot.32–34

Expression of Mouse PIP4K2A and -2B in ARPE-19 Cells Whose PI5P4Kα and β Were Knocked Down by shRNA

The full-length cDNA of mouse PIP4K2A or PIP4K2B was subcloned from pCMVSPORT6-P-IPK2A (3672732; Thermo Scientific, Waltham, MA, USA) or pCMVSPORT6-P-IPK2B (6491155; Thermo Scientific) into a retroviral vector of pLNCX3 (LTR—CMV-SalI-HpaI-SacI-NotI—BgeII—CMV—SAL-I/Not-I). This vector was obtained from Sigma-Aldrich Corp. (St. Louis, MO, USA).

On the day of transfection, plasmid DNA (25 µg) or a transfection reagent (Invitrogen: lipofectamine 2000, 150 µL) was mixed individually with 1.8 mL OptiMEM (Thermo Scientific). The two were then mixed together and allowed to sit at room temperature for 45 minutes. The DNA mixture was added dropwise into approximately 70% confluent HEK 293GPG cells34 with 10 µL OptiMEM in a 25-cm diameter dish. After incubation for 7 to 10 hours, 12 mL virus-producing medium was added, and this day was considered day 0. At 24 hours post transfection, the media were replaced with the complete virus-producing medium. On days 2, 3, 4, and 5, the media were collected into 50mL sterile conical tubes and spun at 800g for 5 minutes. The pooled supernatants were then pelleted in sterile tubes at 25,000g for 90 minutes to concentrate the viruses. Finally, the viral pellets were resuspended in 300 µL sterile TNE buffer (50 mM Tris pH 7.8, 130 mM NaCl, 1 mM ethylenediaminetetraacetic acid [EDTA], transferred into microtubes, and dissolved at 4°C with gentle
Prevention of PVR by Suppression of PI5P4Ks

Rotation overnight. These dissolved retroviruses were then used to infect target cells with 8 μg/mL polybrene or kept at −80°C.32–34 Expression of pLNCX2-PIP4K2A and -2B in the ARPE-19 cells was selected in the 2 mg/mL G418-contained culture medium for 2 weeks and then analyzed by Western blot.

**Western Blot**

Cells grown to 90% confluence in wells of 24-well plates were serum starved for 24 hours, and then treated with or without normal RV (diluted 1:2 in DMEM/F12) for 16 hours. After two washes with ice-cold phosphate-buffered saline (PBS), the cells were lysed in 1× sample buffer diluted with protein extraction buffer [10 mM Tris-HCl, pH 7.4; 5 mM EDTA, 50 mM NaCl, 50 mM NaF, 1% Triton X-100, 20 μg/mL aprotinin, 2 mM Na3VO4, 1 mM phenylmethylsulfonyl fluoride] from 5× sample buffer [25 mM EDTA, 10% sodium dodecyl sulfate (SDS), 500 mM dithiothreitol (DTT), 50% sucrose, 500 mM Tris-HCl (pH = 6.8), 0.5% bromophenol blue]. The samples were boiled for 5 minutes and then centrifuged for 5 minutes at 13,000 g, 4°C. Proteins in the samples were separated by 10% SDS-polyacrylamide gel electrophoresis (PAGE), transferred to polyvinylidene difluoride (PVDF) membranes, and then subjected to Western blot analysis using selected antibodies. Signal intensity was determined by densitometry using NIH ImageJ software.32–34

**Cell Proliferation Assay**

ARPE-19 cells with shRNA for GFP or PIP4K2A/2B were seeded into wells of a 24-well plate at a density of 3 × 10^4 cells/well in DMEM/F12 with 10% FBS. Following attachment, the cells were treated with DMEM/F12 or RV (1:2 dilution in DMEM/F12). On day 3, the cells were trypsin detached from the plates and counted in a hemocytometer. Each experimental condition was assayed in duplicate, and at least three independent experiments were performed.35–37

**Cell Apoptosis Assay**

ARPE-19 cells with shRNA for GFP or PIP4K2A/2B were seeded into 6-cm dishes at a density of 2 × 10^5 cells per dish in DMEM/F12 + 10% FBS. Following attachment, the cells were treated with DMEM/F12 or RV (1:2 dilution in DMEM/F12). On day 3, the cells were stained with fluorescein isothiocyanate (FITC)-conjugated annexin V and propidium iodide (PI) following the manufacturer's instructions (BD Biosciences, Palo Alto, CA, USA). The cells were analyzed by flow cytometry in a Beckman Coulter XI instrument (Brea, CA, USA). At least three independent experiments were performed.32,35

**Cell Migration Assay**

ARPE-19 cells with shRNA for GFP or PIP4K2A/2B were seeded into a 12-well plate were grown to near confluence, and then a wound was created by scratching the monolayer with a 200-μl pipette tip.34 The cells were washed with PBS and treated with DMEM/F12 or RV (1:2 dilution in DMEM/F12). The scratched area was photographed to capture the initial width and photographed again 16 hours later. Analysis was conducted using Adobe Photoshop CS4 software (San Jose, CA, USA). At least three independent experiments were performed.

**Collagen Contraction Assay**

Cells were resuspended in 1.5 mg/mL neutralized collagen I (INAMED, Fremont, CA, USA) (pH 7.2) on ice at a density either of 1 × 10^6 for RPE cells/mL or of 1 × 10^5 for RCF cells/mL.32,35 The mixture was transferred into wells of 24-well plates that had been preincubated overnight with 5 mg/mL BSA/PBS. After the collagen had polymerized at 37°C for 90 minutes, 0.5 mL DMEM/F12 or RV (1:2 dilution in DMEM/F12) was added. On day 3, the gel diameter was measured and the gel area calculated using the formula πr^2, where r is the radius of the gel. At least three independent experiments were performed.35–37

**Experimental PVR in Rabbits**

As previously described,35,36 PVR was induced in the right eyes of Dutch Belted rabbits purchased from Covance (Denver, PA, USA). Briefly, a gas vitrectomy was performed by injecting 0.1 mL perfluoropropane (C8F18) (Alcon, Fort Worth, TX, USA) into the vitreous cavity 4 mm posterior to the corneal limbus. One week later, all rabbits were injected with 0.1 mL platelet-rich plasma (PRP) from rabbits and 0.1 mL DMEM/F12 containing either 3.0 × 10^5 cells of ARPE-19 (10 rabbits) or 1.0 × 10^5 of RCFs (5 rabbits) with shRNA for GFP or PIP4K2A/2B under a surgical microscope. The retinal status was examined with an indirect ophthalmoscope through a +30 D fundus lens on days 1, 3, 5, 7, 14, 21, and 28 by two double-masked ophthalmologists. Proliferative vitreoretinopathy was graded according to the Fastenberg classification from 0 through 5.37 On day 28, animals were killed, the eyes were enucleated, and the eyeballs were either fixed with 10% formalin for histology analysis or frozen at −80°C for vitreous extraction. All surgeries were performed under aseptic conditions and adhered to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. The protocol for the use of animals was approved by the Schepens Animal Care and Use Committee (Boston, Massachusetts).

Rabbit vitreous was prepared from frozen rabbit eyeballs as described previously.34 The left eyes were not manipulated in any way and were normal and free of obvious abnormalities.

**Optical Coherence Tomography (OCT)**

Fundus photographs and OCT were taken using a spectral domain (SD)-OCT system (Bioptigen, Inc., Durham, NC, USA). Rabbits were deeply anesthetized with intramuscular anesthesia consisting of ketamine 30 to 50 mg/kg body weight, xylazine 5 to 10 mg/kg body weight, and acepromazine 1 mg/kg body weight. Depth of anesthesia was verified by the absence of the toe pinch withdrawal reflex. The pupils were dilated with topical 1% tropicamide to view the fundus.38,39

**Immunofluorescence**

Frozen sections of embedded frozen ERMs from three patients with grade C PVR were prepared as described previously,7 and this research adhered to the Tenets of the Declaration of Helsinki and the protocol approved by the Schepens Eye Research Institute Massachusetts Eye and Ear Institutional Review Board (632172-3). The sections were fixed in 3.7% formaldehyde/PBS for 10 minutes and then treated for 15 minutes with 0.3% hydrogen peroxide (to remove endogenous peroxidases). Subsequently, the sections were preincubated with 5% normal goat serum in 0.3% Triton X-100/PBS for 20 minutes, then incubated with primary antibodies against PIP4K2α or PIP4Kβ (1:100 dilution) for 1 hour or with a normal rabbit IgG. After three washes with PBS, the sections were incubated with fluorescently labeled secondary antibodies, Dylight 549 (Vector Laboratories, Inc., Burlingame, CA, USA) (1:300 dilution in a blocking buffer), for 30 minutes. After three washes with PBS, the slides were mounted with a mounting medium containing 4′,6-diamidino-2-phenyldiocol...
previously described, 33 human vitreous (HV) samples from PVR patients (1.0–1.5 mL) were obtained during pars plana vitrectomy performed before initiating the pars plana infusion at the Vancouver Hospital, British Columbia. Ethics approval was obtained before the initiation of this project from the Vancouver Hospital and University of British Columbia Clinical Research Ethics Board. The University of British Columbia Clinical Research Ethics Board policies comply with Tri Council Policy and Good Clinical Practice Guidelines, which have their origins in the ethical principles in the Declaration of Helsinki. Written informed consent was obtained from patients.

Statistics
The data were analyzed using an unpaired t-test or a Mann-Whitney test. A power (p) value less than 0.05 was considered statistically significant.

FIGURE 1. Knockdown of PI5P4Kα and β in ARPE-19 cells led to the suppression of vitreous-stimulated proliferation and survival. (A) ARPE-19 cells expressing shRNA for GFP (GFP) or PI5P4K2A/B (2A/B) were treated for 16 hours with or without rabbit vitreous (RV) diluted (1:2) in DMEM/F12. The cell lysates were subjected to Western blot analysis using the indicated antibodies. Fold change was calculated by first normalizing PI5P4Kα and β for β-actin and then calculating the ratio of the basal (shGFP) to stimulated protein levels for each cell type. This is representative of three independent experiments. (B) The characterized ARPE-19 cells in (A) were seeded into wells of a 24-well plate at a density of 3 × 10⁴ cells/well in DMEM/F12 plus 10% FBS. Following cell attachment, the medium was replaced with 0.5 mL DMEM/F12 or RV diluted (1:2) in DMEM/F12. The cells were counted with a hemocytometer on day 3. The mean ± standard deviation (SD) of three independent experiments is shown; *P < 0.05 using an unpaired t-test. (C) The characterized ARPE-19 cells in (A) were seeded into 6-cm dishes in DMEM/F12 ± 10% FBS at a density of 2 × 10⁵ cells per dish. After 8 hours, the cells were treated with or without RV. On day 3, the cells were analyzed for apoptosis by FACS (fluorescence-activated cell sorting). An example of the raw FACS data is shown. PI, propidium iodide; FITC, fluorescein isothiocyanate conjugated with annexin V for sorting. An example of the raw FACS data is shown. PI, propidium iodide; FITC, fluorescein isothiocyanate conjugated with annexin V for sorting. An example of the raw FACS data is shown. PI, propidium iodide; FITC, fluorescein isothiocyanate conjugated with annexin V for sorting.

EXPERIMENTAL PROCEDURES
Animals
Experiments described in (Fig. 1C) were performed in accordance with the Association for Research in Vision and Ophthalmology (ARVO) Statement for the Use of Animals in Ophthalmic and Vision Research. The experimental design and procedures were approved by the University of British Columbia Animal Care Committee (Animal Use Protocol A09-0533). The test system used was the rabbit cornea. The rabbits were obtained from a local supplier and were housed in a temperature-controlled environment with 12-h light-dark cycles. The rabbit corneas were obtained from rabbits after these animals were assessed and confirmed to be free of any clinical signs of disease. The eyes were enucleated 30 minutes after death and the corneas were removed and placed on a 37°C, humidified incubator with 100% relative humidity. The anterior chamber angle was incised to expose the iris and ciliary body, and a 28-gauge needle was used to create a 1-mm breach in the iris and ciliary body. The eyes were then pressurized with a mixture of 95% oxygen and 5% carbon dioxide at 90 mm Hg for 72 hours. The eyes were then placed in a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then removed and placed on a 37°C, humidified incubator with 100% relative humidity and were left for 14 days to allow the corneas to become clear. The corneas were then remove...
experiments demonstrate that suppression of PI5P4K \(a\) induced migration and contraction of ARPE-19 cells. These and contraction in the serum-starved cells, but did blunt RV-overexpressed (OV) in ARPE-19 cells whose endogenous PI5P4 were suppressed by shRNAs through introduction of \(pLNCX3\)-retroviral vector (EV) \(\text{PIP4K2A}\) to these cells. The cells with an empty retroviral vector (EV) \(pLNCX3\) served as controls. These cells were then treated for 16 hours with or without RV, and their lysates were subjected to Western blot analysis using the antibodies indicated as described in Figure 1A. This is representative of three independent experiments. (B) The characterized ARPE-19 cells in (A) were assessed for their ability to contract as described in the legend of Figure 2B. The mean ± SD of the three independent experiments is shown; * \(P < 0.05\) using an unpaired \(t\)-test, and NS indicates no significant difference between the groups. A photograph of a representative experiment is shown at the bottom of the bar graphs.

results demonstrate that suppression of PI5P4Kz and \(\beta\) attenuates RV-induced cell proliferation and survival against apoptosis, processes intrinsic to the pathogenesis of PVR.\(^{35}\)

**Knockdown of PI5P4Ks in ARPE-19 Cells Blocks Vitreous-Stimulated Cell Migration and Contraction**

PI5P4Kz and \(\beta\) can convert PI-5-P to PI-4,5-P2, which regulates actin filament elongation and interaction of the plasma membrane with the cytoskeleton.\(^{41-43}\) Next, we asked whether knockdown of PI5P4Kz and \(\beta\) impacted RV-induced cell migration and contraction. As shown in Figure 2, suppression of PI5P4Kz and \(\beta\) did not affect cell migration and contraction in the serum-starved cells, but did blunt RV-induced migration and contraction of ARPE-19 cells. These experiments demonstrate that suppression of PI5P4Kz and \(\beta\) blunts RV-induced increase in cellular responses intrinsic to the pathogenesis of PVR, suggesting that knockdown of PI5P4Kz and \(\beta\) might be able to inhibit the development of PVR.

**Expression of Mouse PIP4K2A/2B Restores the Ability of RV-Induced Proliferation and Contraction of PI5P4Kz- and -\(\beta\)-Deficient ARPE-19 Cells**

To exclude unknown nonspecific targets of shRNAs for human \(\text{PIP4K2A/2B}\), we expressed mouse \(\text{PIP4K2A/2B}\) in ARPE-19 cells whose endogenous \(\text{PI5P4Kz}\) and \(\beta\) had been knocked down by shRNAs. As shown in Figure 3A, expression of mouse PI5P4Kz and \(\beta\) was comparable to RV-induced expression of PI5P4Kz and \(\beta\) in ARPE-19 cells. Importantly, expressing mouse PI5P4Kz and \(\beta\) restored the capability of RV-induced contraction (Fig. 3B) of the PI5P4Kz- and \(\beta\)-deficient cells. These experiments indicate that RV-induced expression of PI5P4Kz and \(\beta\) coordinates with p53 loss to promote cell contraction and that PI5P4Kz and \(\beta\) constitute a potential therapeutic target for treatment of PVR.

![Figure 3](image3.png)

**Figure 3.** Expression of mouse PIP4K2A/2B restored RV-induced contraction of ARPE-19 cells whose endogenous PI5P4z and \(\beta\) were suppressed by shRNAs. (A) Mouse PIP4K2A and PIP4K2B (2A/B) were overexpressed (OV) in ARPE-19 cells whose endogenous PI5P4z and \(\beta\) were suppressed by shRNAs through introduction of \(pLNCX3\)-PIP4K2A/PIP4K2B into these cells. The cells with an empty retroviral vector (EV) \(pLNCX3\) served as controls. These cells were then treated for 16 hours with or without RV, and their lysates were subjected to Western blot analysis using the antibodies indicated as described in Figure 1A. This is representative of three independent experiments. (B) The characterized ARPE-19 cells in (A) were assessed for their ability to contract as described in the legend of Figure 2B. The mean ± SD of the three independent experiments is shown; * \(P < 0.05\) using an unpaired \(t\)-test, and NS indicates no significant difference between the groups. A photograph of a representative experiment is shown at the bottom of the bar graphs.

![Figure 4](image4.png)

**Figure 4.** Suppression of PI5P4Kz and \(\beta\) in ARPE-19 cells prevented experimental PVR. (A) PVR was induced in the right eyes of Dutch Belted rabbits as previously described.\(^{33,36}\) Briefly, 1 week after gas vitrectomy, rabbits were intravitreally injected with 0.1 mL PRP and ARPE-19 cells (\(3 \times 10^5\)) expressing shRNA for GFP (GFP) or PIP4K2A/B in 0.1 mL DMEM/F12 (2A/B). The rabbits were examined on days 1, 3, 5, 7, 14, 21, and 28 with an indirect ophthalmoscope. The PVR status for each rabbit was plotted on days 21 and 28. The data were subjected to Mann-Whitney analysis. *A significant difference between the two groups (GFP and -2A/B). (B) Rabbits with PVR representative of stages 0, 2, and 4 were imaged by OCT. Arrows in 0, 2, and 4 point to a normal, a fibrotic, and a pathologic retina with fibrosis banding, respectively. The numbers 0, 2, and 4 at the bottom of the bar graphs represent stages 0, 2, and 4, respectively. (C) Eyeballs from rabbits with PVR stages 0, 2, and 4 were fixed, sectioned, and stained with hematoxylin and eosin. The retina section with PVR stage 4, has been enlarged to in the bottom right to show that there was growth of fibrosis in the abnormal, thicker retina.
Suppression of PI5P4Ks and -β in RCF Cells Abrogates Experimental PVR

Our previous observations demonstrated that RCFs induce an experimental PVR in rabbits that forms more rapidly and is more severe than that induced by ARPE-19 cells. To examine if knockdown of PI5P4Kα and -β could inhibit PVR in rabbits induced by RCFs, the expression of PI5P4Kα and -β was suppressed in RCFs by shRNA (Fig. 5A). Rabbit vitreous also contained PI5P4Kα and -β, as well as a decrease in p53 in these fibroblasts (Fig. 5A). Moreover, suppression of PI5P4Kα and -β reduced RCF contractability that was induced by RV (Fig. 5B). Therefore, we injected these cells and their shGFP control cells intravitreally into rabbit eyes to compare their PVR potential. As shown in Figure 5C, shGFP RCFs led to development of severe PVR (stage 4 or 5). Typically in these stages, retinas get detached totally or close to this degree by day 7, whereas in this study, no rabbits that had been injected with shPI5P4K2A/B RCFs had developed any stage of PVR by day 7 (Fig. 5C). On day 28, the retinas of all rabbits injected with shGFP RCFs had completely detached retinas, while 60% of rabbits injected with shPI5P4K2A/B RCFs remained at stage 0 and 40% had progressed to stage 5 (Figs. 5C, 5D). These experiments indicate that inhibition of PI5P4Kα and -β can prevent severe PVR induced by intravitreal injection of RCFs in rabbits.

Knockdown of PI5P4Kα and -β in RPEM Cells Blocks HV-Induced Contraction

It is believed that RPE cells in ERMs contribute to the pathogenesis of PVR. Thus, we knocked down PI5P4Kα and -β in primary RPE cells isolated from an ERM of a PVR patient (RPEM) (Fig. 6A). Intriguingly, HV also enhanced expression of PI5P4Kα and -β, and reduced p53 levels (Fig. 6A). These cells then were examined for their capability for HV-stimulated contraction in a collagen contraction assay. As shown in Figure 6B, suppression of PI5P4Kα and -β in RPEM cells blocked HV-induced contraction.

The above in vitro and in vivo observations indicate that vitreous-induced expression of PI5P4Kα and -β contributes to the pathogenesis of PVR. Thus, we wondered whether these two proteins were expressed in ERMs from PVR patients. As seen in Figure 6C, immunofluorescence staining revealed that both PI5P4Kα and -β were abundant in the tested ERMs from...
grade C PVR patients and that they were localized to the membranes in virtually all the cells.

**DISCUSSION**

Phosphatidylinositol signaling has been shown to impact a variety of fundamental cellular processes, including intracellular membrane trafficking, cytoskeletal rearrangement, and cell proliferation, survival, and growth.14 In this article, we demonstrate that vitreous stimulates a decrease in p53 and an increase in PI5P4Kz and -b, and that suppression of PI5P4Kz and -b impacted vitreous-induced cell proliferation, survival, migration, and contraction, as well as experimental PVR. It has been reported that depletion of PI5P4Kz and -b inhibits growth of p53-null tumors, where it was suggested that loss of both p53 and PI5P4Kz and -b leads to production of excess reactive oxygen species (ROS) that can lead to reduced cell proliferation and increased senescence.20 Furthermore, the authors speculated that PI5P4Kz and -b required to maintain glucose metabolism, thereby reducing oxidative phosphorylation and enhancing cell proliferation and survival when p53 is deficient.20 We also speculated that a similar mechanism might be underlying the observed effects in PVR. Treatment of RPEs with vitreous leads to the activation of PDGFR/Pi3K/Akt/MDM2, resulting in p53 reduction, and, concurrently, to the enhancement of PI5P4Kz and -b expression and, in turn, to suppress the production of excess ROS, triggering cellular responses intrinsic to PVR.

In addition, PI5P4Kz has been identified as essential for the clonogenic and leukemia-initiating potential of human acute leukemia cells, and it is required for these cells’ proliferation and survival.15 Knockdown of PI5P4Kz in human acute leukemia cells results in accumulation of the cyclin-dependent kinase inhibitors (CDKNs) p21 and p27, G1 cell cycle arrest, and apoptosis.15 However, knockdown of PI5P4Kz in normal hematopoietic stem and progenitor cells does not adversely impact either clonogenic or multilineage differentiation potential.15 Similarly, we found that suppression of both PI5P4Kz and -b did not affect RPE cell proliferation and contraction, but did influence vitreous-induced RPE cellular responses. These findings indicate a differential dependency and suggest that they might be a consequence of the regulation of different transcriptional programs in normal versus RV-induced conditions.

Both PI5P4Kz and -b are capable of phosphorylating PI-5-P on the fourth hydroxyl of the inositol ring to generate PI-4,5-P3, which acts as a substrate for PI3K or phospholipase C to generate the essential second messengers PI-3,4,5-P3 and IP3, respectively.27,41,43,45,46 In addition, PI-4,5-P2 can regulate actin filament elongation, interaction of the plasma membrane with the cytoskeleton, phagocytosis, clathrin-mediated endocytosis, and exocytosis.41,43,47,49 Vitreous-induced PI5P4Kz and -b that leads to increased migration and contraction in RPE cells may be mediated by these biochemical functions. However, it has been reported that knocking down PI5P4Ks activates the PI3K/Akt pathway in a subset of cancer cells, suggesting that the PI5P4Ks suppress PI3K/Akt signaling because they provide an alternative mechanism for enhancing glucose metabolism in response to ROS.26

In this paper, we report that vitreous increased expression of both PI5P4Kz and -b. In addition, both PI5P4Kz and -b were abundant in ERMs from PVR patients. The fact that suppression of PI5P4Kz and -b inhibited only vitreous-induced cellular responses raises the possibility that inhibiting PI5P4Kz and -b might target only cells that have migrated into the vitreous. However, in this study, we used a cell injection rabbit model in which cell proliferation is not from endogenous cell proliferation. To better simulate PVR in clinics, a penetrating retinal injury rabbit model might be a way to test if a potential PI5P4Kz inhibitor, such as tyrphostin AG-82,44 would inhibit ocular trauma-induced PVR. Moreover, these experimental data might provide an option for the future therapeutic targeting of PI5P4Kz and -b to prevent PVR.

**Acknowledgments**

The authors thank Randy Huang for assistance in flow cytometry, Jessica Hoadley and Marie Ortega for their support with rabbit experiments, Oscar Morales for his assistance with OCT, Bianai Fan for histologic sections, and Gale Unger for copyediting this article.

Supported in full by National Institutes of Health (NIH) R01 EY012509 (HL) and in part by NIH National Eye Institute Core Grant P30 EY003790.

Disclosure: G. Ma, None; Y. Duan, None; X. Huang, None; C.X. Qian, None; Y. Chec, None; S. Mukai, None; J. Cui, None; A. Samad, None; J.A. Matsubara, None; A. Kazlauskas, None; P.A. D'Amore, None; S. Gu, None; H. Lei, None

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Prevention of PVR by Suppression of PISP4Ks


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