RKI-1447, a Rho-kinase inhibitor causes ocular hypotension, actin stress fiber disruption and increased phagocytosis

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Abstract

**Purpose:** This study investigated the hypotensive effect of RKI-1447, a Rho kinase inhibitor, in a porcine ex vivo pigmentary glaucoma model.

**Methods:** Twenty-eight porcine anterior chambers were perfused with medium supplemented with $1.67 \times 10^7$ pigment particles/ml for 48 hours before treatment with RKI-1447 (n=16) or vehicle control (n=12). Intraocular pressure (IOP) was recorded and outflow facility was calculated. Primary trabecular meshwork cells were exposed to RKI-1447 or vehicle control; effects on the cytoskeleton, motility, and phagocytosis were evaluated.

**Result:** Compared to baseline, the perfusion of pigment caused a significant increase in IOP in the RKI-1447 group ($P=0.003$) at 48 hours. Subsequent treatment with RKI-1447 significantly reduced IOP from 20.14+/−2.59 mmHg to 13.38+/−0.91 mmHg ($P=0.02$). Pigment perfusion reduced the outflow facility from 0.27+/−0.03 at baseline to 0.18+/−0.02 at 48 hours ($P<0.001$). This was partially reversed with RKI-1447. RKI-1447 caused no apparent histological changes in the micro- or macroscopic TM appearance. RKI-1447-treated primary TM cells showed significant disruption of the actin cytoskeleton both in the presence and absence of pigment ($P<0.001$) but no effect on TM migration was observed. Pigment-treated TM cells exhibited a reduction in TM phagocytosis, which RKI-1447 reversed.

**Conclusion:** RKI-1447 significantly reduces IOP by disrupting TM stress fibers and increasing TM phagocytosis. These features may make it useful for the treatment of secondary glaucomas with an increased phagocytic load.

**Keywords:** Pigmentary glaucoma, ROCK kinase inhibitor, RKI-1447, Trabecular meshwork, Phagocytosis, Cytoskeleton.
Introduction

Ocular hypertension frequently observed in glaucoma patients exhibits a complex pathomechanism that remains incompletely understood [1]. Intraocular pressure (IOP) remains the single most important modifiable variable in glaucoma treatment [2]. Current topical medications for glaucoma typically increase the uveoscleral outflow of aqueous humor (prostaglandin analogs) or reduce aqueous humor production (beta-adrenergic blockers, alpha-adrenergic agonists, carbonic anhydrase inhibitors). Until recently, miotics were the only class that increases the conventional outflow as the primary mechanism but were hampered by significant side effects that include ciliary spasm and dim vision caused by miosis [2]. Newly introduced nitric oxide (NO) donors increase the facility by relaxing the TM [3] and the distal outflow tract [4, 5]. Rho-kinase inhibitors also relax the TM cytoskeleton [6, 7]. In the present study, we examine the hypotensive effect of RKI-1447, which was originally introduced as an anti-tumor agent in breast cancer treatment [8].

Rho-associated protein kinase (ROCK) is part of the serine-threonine kinases that act on the cytoskeleton to regulate the movement and shape of cells. It plays a central role in biological functions including stress-fiber formation and contraction, cell adhesion, migration and invasion, transformation, phagocytosis and apoptosis, cytokinesis and mitosis as well as differentiation [9, 10]. Previous studies found that ROCK is expressed in the TM and upregulated in the optic nerve head of glaucomatous eyes [11, 12]. Honjo and colleagues found the ROCK inhibitor Y-27632 increased outflow facility in a dose-dependent manner in rabbit eyes [13]. The investigators suggested that the hypotensive effect may result from a disruption of actin bundles and reduction in focal adhesion of TM cells [13]. Similarly, recent studies found that ROCK inhibitors disrupt tight junctions thereby increasing the permeability of Schlemm’s canal endothelium [14, 15] in addition to decreasing the episcleral venous pressure [16].

Ripasudil (K-115) was the first approved ROCK inhibitor for glaucoma [6, 7]. Two randomized trials involving 413 Japanese primary open-angle glaucoma patients found that Ripasudil achieved an average of 0.4-1.6 mmHg larger IOP reduction than the administration of timolol or latanoprost alone [17]. Phase II/III trials that aim to investigate the IOP-lowering effect of four different ROCK inhibitors (AMA0076 [18], AR-13324 [19], K-115 [7] and PG-324 [20]) are currently in process. Several trials that employed functionally and structurally similar compounds were discontinued due to the low therapeutic efficiency and high incidence of adverse events [21].

First characterized by Patel et al. in 2012 [8], RKI-1447 inhibits the type I kinase by binding to the ATP binding site through interaction with the Asp-Phe-Gly motif and hinge region (Figure 1) [8, 22]. In vitro, it selectively inhibits ROCK1/2-mediated actin stress fiber formation at a low IC50 (14.5 nM and 6.2 nM for ROCK1 and ROCK2, respectively [23]) following lysophosphatidic acid stimulation. Unlike other ROCK inhibitors, RKI-1447 does not exhibit cross-reactivity with AKT, MEK, PKA, and PKG signaling, even at a high concentration of up to 10 μM [8, 23], reducing the risk of side effects. Recently, our group developed an ex vivo pigmentary glaucoma model that exhibited IOP elevation, typical TM stress fiber formation, and reduction in TM phagocytosis [24, 25]. In this paper, we examine the ocular hypotensive effect of RKI-1447 in an ex vivo anterior chamber perfusion model and investigate its possible mechanisms.
Materials and Methods

Pig eye perfusion, pigmentary glaucoma model, and RKI-1447 intervention

Because eyes were obtained from an abattoir, no Institutional Animal Care and Use Committee protocol was required. “Porcine anterior segment perfusion culture is a well-established homolog to human and primate anterior segment culture, with porcine outflow tracts being similar in anatomy and more readily attainable within short time-periods after death [26, 27].” The setup of the porcine eye perfusion system followed our previous protocols [24, 28–30]. Briefly, 40 freshly enucleated pig eyes were obtained from a local slaughterhouse (Thoma Meat Market, Saxonburg, PA) and processed within two hours of sacrifice. After the removal of extraocular tissues, the eyes were disinfected in a 5% povidone-iodine ophthalmic solution (betadine 5%; Alcon, Fort Worth, TX, USA) for two minutes, washed three times with PBS, and hemisected. Anterior chambers were then mounted on the perfusion dishes and subsequently underwent continuous perfusion with 1% FBS supplemented with DMEM and antibiotics at a constant rate of 3μl per min.

Once baseline IOPs were obtained, 28 of the 40 eyes were perfused with medium supplemented with \(1.67 \times 10^7\) pigment particles/ml for 48 hours before being subjected to the RKI-1447 (n=16) or the vehicle control (n=12). For the RKI-1447 group, RKI-1447 was diluted into the pigment medium to a final concentration of 1μM. For the vehicle control, eyes were continuously perfused with the same pigment medium. A total of 12 of the 40 samples without pigment perfusion and RKI-1447 served as the normal control. IOPs were measured using a pressure transducer at two-minute intervals.

Production of pigment dispersion particles

Pigment dispersion particles were produced as previously described with a freeze-thaw protocol [31, 32]. In brief, ten porcine irises were sealed in a 50-ml tube, frozen at a temperature of -80°C for 2 hours, then thawed at room temperature. After two cycles of freeze-thaw, 15 ml of phosphate-buffered saline (PBS) was added to the tube, and the mixture was pipetted up and down 100 times with a 3 ml Pasteur pipette before straining through a 70-μm cell strainer (cat#431751, Corning Incorporated, Durham, NC, USA). The suspension was centrifuged at 3000 rpm for 15 min. The supernatant was discarded, and the pigment pellet was resuspended in PBS and centrifuged again at the same settings. Finally, pigment particles were resuspended in 4 ml of PBS as the stock solution and titered using a hemocytometer (#1490, Hausser Scientific, Horsham, PA) at 1000-fold dilution as per the method previously described [32].

Histology

At the conclusion of the experiments, the anterior segments were dismounted, washed with PBS, fixed with 4% PFA for 24 hours and embedded in paraffin. Samples were sectioned at 6-micron thickness, followed by staining with hematoxylin and eosin (H&E).

Primary TM culture

Primary porcine TM cells were isolated and characterized as described before [32]. A porcine anterior segment was dissected by removing the sclera and cornea and creating TM strips of 1 mm. Three pieces were cultured in a T75 flask with TM culture media containing 5% FBS supplemented with Opti MEM (31985-070, Gibco, Life Technologies, Grand Island, NY) and antibiotics (15240062, Thermo Fisher Scientific, Waltham, MA). The TM cells had migrated onto the bottom of the flask by day four and formed clones. Cells were passaged at 80% confluence and replated at a 1:3 ratio. We characterized these cells by immunostaining with a combination of TM specific antibodies (alpha smooth muscle actin [alpha-SMA] antibody, matrix gla protein...
[MGP] antibody, and aquaporin 1 [AQP1]), an in vitro phagocytic assay and myocilin induction by glucocorticosteroids [32]. To avoid transformation, differentiation, and senescence, only cells with up to four passages were used.

**Immunostaining**

For further characterization, we plated the TM cells onto 10 X 10 mm glass slides and fixed them with 4% PFA at 80% confluence. This was followed by washing with PBS three times and incubation with the primary antibody overnight at 4°C. Primary antibodies were goat polyclonal anti-MGP antibodies (1:100 dilution, sc-32820, Santa Cruz, Dallas, Texas), rabbit polyclonal anti alpha-SMA (1:100, ab5694, Abcam, Cambridge, MA) and rabbit polyclonal anti-AQP1 antibodies (1:100, Sc-20810, Santa Cruz, Dallas, Texas). After three rinses in PBS, a mixture of secondary antibodies (donkey-anti-goat Alexa Fluor® 647 [1: 200, ab150131, Abcam, Cambridge, MA] and goat anti-rabbit Alexa Fluor® 488 [1:1000, A27034, Thermo Scientific, Waltham, MA]) was incubated with the cells for 45 minutes at room temperature. Cell nuclei were counterstained with DAPI (1: 1000, D1306, Thermo Fisher Scientific, Waltham, MA) for 15 minutes. Images were taken using an upright laser scanning confocal microscope at 400x magnification (BX61, Olympus, Tokyo, Japan).

To quantify the stress fiber formation, cells were fixed with 4% PFA for 24 hours before being stained with Alexa Fluor® 488 Phalloidin for 30 minutes (1:20, A12379, Thermo Scientific, Waltham, MA). Cell nuclei were labeled with 4',6-diamidino-2-phenylindole (DAPI). Ten randomized visual fields were chosen to quantify the percentages of cells with stress fibers for each group.

**TM cell motility assay**

Evaluation of in vitro TM cell motility utilized a scratch assay with a minor modification[33]. In brief, 2×10⁵ primary TM cells were seeded into each well of a six-well plate and cultured in an incubated chamber equipped with a digital camera and image acquisition software. After reaching 80% confluence cells were treated with pigment particles at a concentration of 1.67×10⁷ particles/ml for 24 hours. A 10μl pipette tip (info) was used to create a cell-free scratch that had a consistent width. RKI-1447 (final concentration: 1μM) or vehicle was applied. Pictures from ten positions of each well were acquired at 40-fold magnification in two-hour intervals for 20 hours. The numbers of TM cells which moved into the cell-free scratch were counted.

**TM cell phagocytosis assay**

Phagocytic activity of the primary TM cells was measured by flow cytometry using a fluorescent microsphere ingestion test [29]. 2×10⁵ primary TM cells were seeded into each well of a six-well plate and exposed to 1.67×10⁷ pigment particles/ml or vehicle treatment for seven days. Half of the cells were then treated with 1μM RKI-1447 for 24 hours while the rest underwent sham treatment. To quantify the phagocytic activity, cells were incubated with 5×10⁸ 0.5-micron fluorescent microspheres per milliliter at 37°C for 1 hour. After washing thoroughly with PBS, cells were dissociated with trypsin and resuspended in 200μl PBS. The percentage of cells which had ingested microspheres was determined by flow cytometry.

**Statistics**

Quantitative data were presented as the mean +/- standard error and processed by PASW 18.0 (SPSS Inc., Chicago, IL, USA). A paired t-test was used to compare IOP changes from baseline. Other quantitative data were compared by one-way ANOVA. A p≤0.05 was considered statistically significant.
Results

IOP reduction and increase of outflow facility

Baseline IOPs among these three groups were comparable (10.32 +/- 0.82 mmHg in the pigment group versus 12.21 +/- 1.06 mmHg in the RKI-1447 group and 10.21 +/- 1.15 mmHg in the control group, P=0.291). Compared to each baseline, the intracameral perfusion of pigment particles resulted in a significant IOP elevation in the pigment group (P=0.038) and the RKI-1447 group (P=0.003) at 48 hours, while IOPs in the control group remained unchanged (P=0.108). RKI-1447 significantly reduced IOP from 20.14 +/- 2.59 mmHg to 13.38 +/- 0.91 mmHg at 60 hours and remained at this low level throughout the study. In contrast, IOP in the pigment group remained high in comparison to its baseline (all P<0.05). At all time points after RKI-1447, IOP in the RKI-1447 group exhibited no statistical difference compared to the control (all P>0.05); however, both were significantly lower than those in the pigment group (all P<0.05) (Figure 2A).

Because the episcleral venous pressure is close to zero in the perfusion system, outflow facility is inversely correlated with IOP. In the current study, baseline outflow facility exhibited no difference across the three groups (P=0.316). Pigment perfusion caused a significant reduction in outflow facility in the RKI-1447 group, from 0.27 +/- 0.03 at the baseline to 0.18 +/- 0.02 at 48 hours (P<0.001), while RKI-1447 partially reversed this effect by 14.57% to 48.35% at the follow-up time points. In contrast, outflow facility in the pigment group continuously decreased from a baseline of 0.31 +/- 0.03 to 0.15 +/- 0.01 at the end of the study (all P<0.05, compared to its baseline). In the control group, outflow facility exhibited a slight decrease as time elapsed (Fig. 2B).

Normal TM exhibits a light-pigmented, multilayer, strainer-like structure connected with the aqueous plexus (Figure 3a). Consistent with our previous findings[24, 29], pigment perfusion resulted in an increased pigmentation in the TM, while most of the pigment particles seemed to be phagocytosed by TM cells without physically obstructing the TM (Figure 3b). RKI-1447 treatment exhibited no apparent changes in TM histology (Figure 3c).

Disruption of TM cell actin stress fibers

The primary TM cells had an elongated morphology (Figure 4a), active phagocytosis (Figure 4b), and were positive for TM markers alpha-SMA, AQP1, and MGP (Figures 4c and 4d). Pigment dispersion did not cause an apparent morphological change in TM cells [29] (Figures 4e and 4g), while RKI-1447 stimulated a cell body contraction within minutes, in the presence or absence of pigment exposure (Figures 4f and 4h).

F-actin cytoskeleton labeling with Alexa Fluor® 488 Phalloidin, showed a baseline of 32.57 +/- 2.66 % actin stress fibers in normal TM cells, compared to 67.13 +/- 4.00 % in the pigment-exposed TM cells (P<0.001) (Figures 4i and 4k). RKI-1447 abolished these stress fibers, both in normal TM cells (2.57 +/- 0.86 % versus 32.57 +/- 2.66 %, P<0.001) and pigment-exposed TM cells (2.51 +/- 0.71 % versus 67.13 +/- 4.00 %, P<0.001) (Figures 4j, 4i, and 4m).
TM cell motility

Because TM cells of glaucomatous eyes [29, 34] or TM cells exposed to pigment dispersion [32], have altered motility, we tested the effect of RKI-1447 on motility using a modified scratch assay. As shown in Figure 5, the pigment exposure statistically decreased the numbers of migrating TM cells (60.83±3.46 in the normal control versus 31.75±3.36 cells in the pigment treated group, P<0.001) while RKI-1447 seemed to have no effect on the TM migration, either for the normal TM cells or the pigment-treated TM cells (P=0.091 and P=0.398, compared to each baseline).

As we have seen before [24, 32], pigment exposure caused a reduction in TM phagocytosis by 10.31% while RKI-1447 reversed this effect and increased it to 111.52% in comparison to the normal baseline.

Discussion

Pigmentary glaucoma (PG) is a common type of secondary open-angle glaucoma with an incidence of 1.4 per 100,000 [35, 36]. A readily recognizable feature of this disease is the dispersion of pigment onto intraocular structures, especially on the corneal endothelium and the trabecular meshwork [37]. Our previous study in a porcine eye perfusion model suggested that iris pigment particles induced TM stress fibers, reduced TM phagocytosis, adhesion, and motility, and caused an IOP elevation [29]. In the current study, we found that RKI-1447, a ROCK1/ROCK2 inhibitor, efficiently reduced the IOP by 33.58% and increased the outflow facility by 38.25% in the same pigmentary glaucoma model. The in vitro results suggest these changes result from an extensive disruption of TM actin stress fibers. In the current study, we utilized a porcine anterior segment perfusion model that mimics the TM changes and IOP phenotype of human PG to examine the hypotensive effect of RKI-1447, a drug previously used in breast cancer treatment [8]. Pigment dispersion caused a significant IOP elevation that plateaued at 48 hrs, and this was partially reversed by RKI-1447. Cytoskeletal changes of TM cells, their stiffness, adhesion, migration, contraction, and phagocytosis can all impact the outflow facility [38, 39], and ROCK signaling plays a central role in all these [11, 40].

The impact of the actin cytoskeleton and cell or extracellular matrix stiffness on aqueous outflow facility has been well established [41, 42]. Consistent with our previous work [29], only a small portion of stress fibers formed in normal control TM while pigment exposure caused a 2.06-fold increase. Actin stress fibers contribute to the stiffness of both the TM cells and the extracellular matrix that surrounds them [41, 43, 44] which directly impact aqueous outflow facility [42]. Gavara et al. also found that the presence of aligned and thick stress fibers in the cell periphery gave rise to reinforced cell stiffness [45]. In general, factors that induce stress fiber formation increase TM stiffness and decrease outflow facility as is the case with senescence and aging, risk factors for primary open-angle glaucoma [46]. In a different study, dexamethasone, an inducer of cross-linked actin networks [47, 48], was shown to increase in three days the stiffness of primary TM cells twofold and that of extracellular fourfold [44]. Clark et al. observed an IOP elevation in ex vivo perfused human anterior segments [48]. In contrast, several reagents that disrupt actin stress fibers exhibit an ocular hypotensive effect. For instance, Y27632 has been found to abolish cyclic stress-induced stress fiber and increase the facility by 40%-80% in pig eyes [49]. Other molecules that target the myosin light chain and its kinase exhibited a similar effect [50–52]. In the current study, RKI-1447 significantly disrupted stress fibers in pigment-treated TM cells, and the IOP-reducing effect could be at least partially attributed to this.
As is the case with other ROCK inhibitors [13, 52, 53], we found RKI-1447 caused TM cell contraction, rounding and decreased adhesion. Live cell imaging indicated that pigment exposure reduces TM cell migration, but this was unaffected by RKI-1447, suggesting that the IOP-lowering effect might not be directly related to TM motility.

Phagocytosis requires a well-coordinated rearrangement of the actin cytoskeleton [54, 55]. TM cells can phagocytose debris to prevent it from entering the outflow tract [38, 56]. A reduction in TM phagocytosis has been observed in different types of glaucoma and glaucoma models. For instance, perfusion of human anterior segments with dexamethasone for 21 days resulted in a 57% decrease of TM phagocytosis without TM cell loss [57]. Switching B-Cell Complex 70 kDa Subunit (SWAP70), vasodilator-stimulated phosphoprotein (VASP), and Wiskott–Aldrich Syndrome protein (WASp), proteins involved in actin polymerization and phagocytosis, were believed to play an essential role in this process [32, 58]. Our observation here of a 10.3% reduction in TM phagocytosis following pigment exposure and increase an 11.5% increase by RKI-1447 matches this.

Our study had several limitations. The anatomical difference between the human and the porcine outflow tract and their angular aqueous plexus is significant [59]. Domestic pigs, even breeders with a longer lifespan, do not typically develop glaucoma. A study by Camras et al. suggested that TM stiffness was correlated with outflow facility in human eye perfusion but not in the pig specimens [60]. This ex vivo study does not account for critical factors that might only be detectable in vivo. It is possible that RKI-1447 does not only have an impact on the TM but also causes a reduction of post-trabecular outflow resistance [4, 5].

In conclusion, the ROCK inhibitor RKI-1447 disrupted TM stress fibers, increased phagocytosis and lowered IOP in an ex vivo model of pigmentary glaucoma.

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Conflict of interest
The authors declare that they have no conflict of interest.
References


Figure 1. Molecular structure of RKI-1447. (a) RKI-1447 is a cell-permeable pyridyl thiazolyl-urea that acts as an ATP site-targeting Rho kinase inhibitor with an IC50 of 14.5 nM and 6.2 nM for ROCK1 and ROCK2, respectively. (b) RKI-1447 interacts with the Asp-Phe-Gly motif and hinge region of ROCKs (blue).
Figure 2. Changes in intraocular pressure and outflow facility by RKI-1447 treatment. (A) IOP at different time points is shown. RKI-1447: blue diamonds, normal control: orange Xs, pigment: black triangles. Measurements were taken 48 hours after pigment-induced, stable IOP elevation. After treatment with RKI-1447, IOP of the RKI-1447 group decreased significantly from 20.14 ±2.59 mmHg to 13.38±0.91, while the pigment group remained significantly higher than its baseline (all P<0.05). (B) The reduced facility could be increased by RKI-1447. In the pigment group, outflow facility was significantly reduced at all time points in comparison to baseline (all P<0.05).
Figure 3. TM histology. The TM in the control group presented as a lightly-pigmented strainer-like tissue, consisting of reticular lamella with TM cells (left). TM of pigment dispersion eyes showed phagocytosed pigment more commonly in the uveal and corneoscleral layers (center: pigment dispersion eye; right: pigment dispersion eye treated with RKI-1447 (right). TM: trabecular meshwork. AP: porcine angular aqueous plexus, with Schlemm’s canal-like segments. CE: corneal endothelium.
Figure 4. Primary TM characterization, cell morphology, and actin cytoskeleton. Primary porcine TM cells were phagocytically active (4a) and tested positive with the preferential TM-marker antibodies, MGP, AQP-1, and alpha-SMA (4b-d). Pigment exposure did not stimulate a significant morphological change (4e, f). Cells contracted following RKI-1447 (g, h). RKI-1447 significantly disrupted the actin cytoskeleton regardless of presence or absence of pigment (P<0.001) (4i-l). Insets in 4i-l correspond to magnified locations in 4m-p. Red arrows: F-actin stress fibers, white arrows: non-bundled actin filaments.
Figure 5. RKI-1447 did not affect TM motility. Motility of TM cells was measured using a scratch assay. Compared to the normal control, pigment exposure resulted in a significant reduction in the number of migrating TM cells (60.83±3.46 in the control versus 31.75±3.36 cells in the pigment group, P<0.001). RKI-1447 had no significant effect on normal control TM cells (P=0.091) or pigment dispersion TM cells (P=0.398).