Purinergic dysregulation causes hypertensive glaucoma–like optic neuropathy

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Introduction

Glaucoma is the second most common cause of blindness and affects more than 70 million people worldwide (1, 2). Elevated intraocular pressure (IOP) is one of the highest risk factors for glaucoma, and lowering IOP is the first-line therapy, which prevents or delays vision loss (3, 4). ATP and nucleotides are present in the aqueous humor (5). Nucleotides are released from various tissues and cells, including lens (6), trabecular meshwork (7), whole retina (8), corneal endothelial cells (9), retinal ganglion cells (RGCs) (10), retinal astrocytes (11), and ciliary body (CB) (12). ATP levels in the aqueous humor are highly elevated in glaucoma patients (13, 14), and although IOP and ATP levels in the aqueous humor are positively correlated in patients (14), their interaction is undefined. Extracellular nucleotides including ATP bind to purinergic P2 receptors (15). P2 receptors are divided into P2X and P2Y receptor subfamilies. Among P2Y receptors, P2Y1 receptor is expressed in cornea, ciliary processes, and trabecular meshwork (16), and the P2Y2 receptor is expressed in cornea, ciliary processes, and retinal pigmented epithelium. P2Y4 and P2Y6 receptors are present in cornea, ciliary processes, photoreceptors, and ganglion cells. P2Y11 receptor is expressed in the retinal pigmented epithelium. Of these, P2Y6 (17, 18), P2Y2 (19), and P2Y11 receptors (20) have been shown to control IOP; however, it is unclear how these receptors control IOP and what the pathophysiological consequence of their dysregulation is. In the present study, we report that the P2Y6 receptor is responsible for IOP reduction and that its dysfunction causes hypertensive glaucoma–like optic neuropathy.
Results

P2Y6 receptor activation reduces IOP. We investigated the physiological role of P2Y6 receptor on IOP. Instillation of uridine diphosphate (UDP) (5 μl/eye, 1.5 hours), an endogenous agonist for P2Y6 receptor, caused IOP reduction of 3-month-old WT mice in a concentration-dependent manner (Figure 1A). At 500 μM, UDP showed a maximum effect of a 10% reduction of IOP. Significant IOP reduction by UDP was obtained at 1.5–6 hours after its application, and the maximum effect was obtained at 1.5 hours after application (Figure 1B). The IOP at 24 hours showed a tendency to be elevated and recovered to the initial level at 48 hours. (C) The effect of an antagonist for P2Y6 receptor on IOP of WT mice. MRS2578 (30 μM, 1.5 h), a selective antagonist for P2Y6 receptors, increased the IOP (n = 10, *P < 0.05, Mann-Whitney U test). (D) The UDP effect in 3-month-old P2Y6 receptor-deficient (P2Y6KO) mice. UDP (500 μM, 1.5 h) did not change the IOP in P2Y6KO mice (n = 20, P = 0.801, Mann-Whitney U test). Data are shown as means ± SEM.

Figure 1. P2Y6 receptor activation reduces intraocular pressure. (A) Concentration dependency of the intraocular pressure (IOP) lowering effect by instillation of uridine diphosphate (UDP) in 3-month-old WT mice. UDP (1.5 h) showed a concentration-dependent IOP lowering effect, and the maximum effect was obtained at 500 μM (n = 20, **P < 0.01 and *P < 0.05 vs. saline-treated eyes, 1-way ANOVA followed by Fisher’s least significant difference [LSD] test). (B) Time dependency of the effect of UDP on WT mice. UDP (500 μM) showed no IOP changes at 30 min but induced a maximum IOP changes at 1.5 hours, followed by gradual recovery. Statistically significant effects were obtained at 1.5–6.0 hours (n = 20, **P < 0.01 and *P < 0.05 vs. initial IOP value, 1-way ANOVA followed by Fisher’s LSD test). The IOP at 24 hours showed a tendency to be elevated and recovered to the initial level at 48 hours. (C) The effect of an antagonist for P2Y6 receptor on IOP of WT mice. MRS2578 (30 μM, 1.5 h), a selective antagonist for P2Y6 receptors, increased the IOP (n = 10, *P < 0.05, Mann-Whitney U test). (D) The UDP effect in 3-month-old P2Y6 receptor-deficient (P2Y6KO) mice. UDP (500 μM, 1.5 h) did not change the IOP in P2Y6KO mice (n = 20, P = 0.801, Mann-Whitney U test). Data are shown as means ± SEM.
or topical application of sodium fluorescein (Supplemental Figure 1B, referred to as Fph-B). In these analyses, a CCD camera was placed above the mouse, and the iris was not dilated (Supplemental Figure 1C). In this arrangement, fluorescent changes in the anterior chamber can be imaged (Figure 3A). Under these conditions, fluorescent levels in the anterior chamber were significantly increased during monitoring for 30 minutes. Although systemically or i.v.-injected fluorescein is usually used for fluorescent angiography, we did not observe dye leakage from retinal vessels (data not shown). The aqueous humor, isolated by piercing the cornea, showed a significant increase in fluorescence levels. The fluorescent intensities of isolated aqueous humor at 5 and 30 minutes were 200% (Supplemental Figure 1D) and 1,500% (Supplemental Figure 1E) of the initial levels, respectively. When we performed time-lapse imaging of WT mice using the Fph-A model, the fluorescence intensity was constantly increased and reached 300% of the initial level at 30 minutes after injection (Figure 3B). Instillation of UDP (5 μl/eye at 500 μM) at 30 minutes before the fluorescein injection significantly suppressed the increase in fluorescence. Timolol (5 μl/eye at 0.5%, 30 min), a β blocker, also suppressed the increase in fluorescence (Figure 3C). Latanoprost (5 μl/eye at 0.005%), a prostaglandin F2α analogue, did not suppress the fluorescence changes in Fph-A (Figure 3, D and E). The speed of fluorescence increase was significantly enhanced in P2Y6KO mice (>400% at 30 min) (Figure 3F). The suppression by UDP disappeared in P2Y6KO mice (Figure 3, G and H). In contrast, the Fph-B model showed an approximate 20% reduction in fluorescence intensity of the anterior chamber during 30-minute monitoring (Supplemental Figure 1, F and G). For Fph-B, latanoprost significantly enhanced the reduction in fluorescence (Figure 3, I and J). In contrast, both UDP and timolol showed no suppressive effects (Figure 3J and Supplemental Figure 1G). We did not find any effect caused by a deficiency of P2Y6 receptor on Fph-B (Figure 3J). These data suggest that P2Y6 receptor activation modifies the dynamics of the aqueous humor in a similar way to timolol. If P2Y6KO mice showed sustained changes in aqueous humor dynamics, their IOP should be changed. As suspected, P2Y6KO mice exhibited elevated IOP compared with WT mice at the ages of 3, 6, 12, and 18 months (Figure 3K). Because the P2Y6 receptor is expressed also in the cornea (16) and many IOP measurement methods for humans have been reported to be affected by central corneal thickness (CCT), we evaluated CCT and structural abnormalities of the cornea. We did not find any structural abnormalities (Supplemental Figure 2A) or thicknesses of the cornea (Supplemental Figure 2B) in mature adult mice (3 months old). In middle-aged mice (12 months old), the CCTs of WT and P2Y6KO mice were both increased (Supplemental Figure 2B). The middle-aged P2Y6KO mice showed a thicker CCT than that of the middle-aged WT mice. No obvious structural abnormalities of the cornea were observed in middle-aged WT or P2Y6KO mice.

P2Y6KO mice show histological abnormalities at middle age. Elevated IOP is one of the highest risk factors for glaucoma (3, 4); therefore, we determined whether P2Y6KO mice had histological abnormalities related to glaucoma. Because axonal damage is one of the first histological changes in glaucoma, we investigated the axonal structure using serial block-face scanning electron microscopy (SBF-SEM) (Figure 4A, asterisks). To obtain detailed ultrastructural information, we collected a large number of serial images of the optic nerve (Figure 4B). SBF-SEM did not show substantial abnormalities, such as axonal swelling or
Figure 3. Aqueous humor dynamics. (A) Fluorophotometry model A (Fph-A). I.p.-injected fluorescein concentrated in the anterior chamber. (B-D) Time courses of fluorescent changes in the anterior chamber. (B) UDP and (C) timolol reduced the fluorescein influx rate, but (D) latanoprost did not ($n=10-21$, **$P<0.01$ vs. control, 2-way repeated ANOVA followed by Fisher’s LSD test). (E) Quantitative data for fluorescence intensity at 30 minutes ($n=10-21$, *$P<0.05$, **$P<0.01$ vs. control, 1-way ANOVA followed by Fisher’s LSD test). (F) P2Y6KO mice showed enhanced fluorescein influx, and (G) no UDP effects.
were observed. (H) Quantitative data at 30 minutes of F and G (n = 10–16, *P < 0.05 vs. control, 1-way ANOVA followed by Fisher’s LSD test). (I) Representative images of latanoprost-treated eyes in the Fph-B model. Significant reduction in fluorescence intensity was observed at 30 minutes. (J) Fluorescein draining was significantly enhanced by latanoprost but not by UDP or timolol, or in P2Y6KO mice (n = 10–18, **P < 0.01 vs. control, 1-way ANOVA followed by Bonferroni correction). (K) IOP levels were significantly higher compared with those of WT mice (n = 61–318, ***P < 0.01, Bonferroni correction). Data are shown as means ± SEM. Scale bars: 2 (A, inserts, and I) and 5 mm (A).

accumulation of axoplasmic organelles in 3-month-old WT, 3-month-old P2Y6KO, and 12-month-old WT mice (Figure 4C). However, the 12-month-old P2Y6KO mice showed prominent axonal swelling with axoplasmic organelle accumulation in the optic nerve (Figure 4C, arrows).

We then analyzed histological changes in the retina using optical coherence tomography (OCT). We estimated the thickness of ganglion cell plus inner plexiform layers (GCL/IPL) because the thinning of these layers is correlated with the loss of visual function in glaucoma patients (23–25) and experimental glaucoma model in primates (26, 27). Tomographic images were obtained from around the optic nerve head (indicated as a green circle in the upper panels of the OCT images in Figure 4D). WT mice showed no significant changes in the thickness of GCL/IPL among 3-, 6-, and 12-month-old mice (Figure 4D). Three-month-old P2Y6KO mice had a similar GCL/IPL thickness with WT mice, but 6- and 12-month-old P2Y6KO mice had a significantly thinner GCL/IPL. The other retinal layers, including the outer plexiform layer (OPL), outer nuclear layer (ONL), external limiting membrane (ELM), and inner/outer segment (IS/OS), showed no structural abnormalities by OCT (Supplemental Figure 3A). We did not find any significant abnormalities in the structure of the CB or anterior chamber (including the trabecular meshwork and Schlemm’s canal) in P2Y6KO mice (Supplemental Figure 3B). Next, we performed the quantitative analysis of RGC damage by immunofluorescence staining. There was no difference in the number of BRN3A+ RGCs in 3- to 12-month-old WT mice (Figure 4E). In 3-month-old P2Y6KO mice, the number of BRN3A+ RGCs was similar to that of WT mice. P2Y6KO mice at 6 and 12 months old showed a significant reduction in RGC numbers.

We then evaluated whether RGC damage in P2Y6KO mice was caused by sustained IOP elevation or by a deficiency of the P2ry6 gene. Lowering IOP using a topical agent prevents or delays vision loss (3, 4); therefore, we used latanoprost to reduce IOP levels in P2Y6KO mice. Daily administration of latanoprost (0.005%, 5 μl/eye/day, from 1–6 months old) reduced IOP levels of P2Y6KO mice to the level of 6-month-old WT mice (Figure 5A). Under this condition, the reduced GCL/IPL thickness (Figure 5B) and RGC numbers (Figure 5, C and D) were restored. Thus, our data show that the sustained elevation of IOP is a cause in P2Y6KO mice.

Impaired visual function in P2Y6KO mice. Because the degeneration of RGCs is a primary cause of blindness in glaucoma, we tested the visual function of P2Y6KO mice using multifocal electroretinograms (mERGs). Four-month-old WT, 4-month-old P2Y6KO, and 13-month-old WT mice showed no significant dysfunction, whereas 13-month-old P2Y6KO mice showed significantly impaired visual function (Figure 6). Taken together, these findings indicate that middle-aged but not mature adult P2Y6KO mice have disordered visual function.

Downregulation of P2Y6 receptors in WT mice with age. We evaluated whether such impairments of the P2Y6 receptor occur naturally in the eye. Because aging is another key risk factor for pathogenesis of glaucoma (28–31), we investigated whether aging affects P2Y6 receptor levels and/or functions. We evaluated retinal P2ry6 mRNA levels and found that retinae from WT mice at 6–12 months old had significantly lower P2ry6 mRNA levels compared with 1- to 3-month-old WT mice (Supplemental Figure 4A). This result was in accord with a previous study (32). Because it was difficult to isolate CB, we estimated the expression level of P2Y6 receptor in the CB by immunofluorescence staining. Fluorescence signals for P2Y6 receptor were detected in the CB (Supplemental Figure 4B), and the signal intensity in 12-month-old WT mice reduced by 50% compared with 3-month-old WT mice. Because the layered pattern of P2Y6 receptor signals correspond with NPE cells, we examined P2Y6 receptor functions by Ca2+ imaging. Isolated ciliary bodies/sclera from mature adult WT mice were sliced by a tissue chopper and plated on a culture dish. Pigmented epithelial cells (PE cells) can be identified by the pigment in their cytoplasm. UDP (100 μM) evoked clear intracellular calcium concentration ([Ca2+]i) transients in NPE cells (Supplemental Figure 4C) but not in PE cells (Supplemental Figure 4D). UDP-evoked [Ca2+]i responses were abolished in the presence of MRS2578 (3 μM) (Supplemental Figure 4E), indicating that functional P2Y6 receptor is expressed in NPE cells. The Ca2+ responses in NPE cells were significantly reduced when they were isolated from 12-month-old WT mice (Supplemental Figure 4, F and G). We determined whether the reduction in P2Y6 receptor–mediated responses contributed to the effectiveness of
UDP-induced IOP lowering. Instillation of UDP (500 μM, 5 μl/eye for 1.5 hours) in 12-month-old WT mice still reduced the IOP (Supplemental Figure 4H), but the efficacy was reduced to about 50% of that in 3-month-old WT mice (4.6% ± 3.0% and 10.1% ± 2.8% for 3- and 12-month-old mice, respectively).

Discussion

We report that the P2Y₆ receptor is essential for regulating the IOP, and deficiency of this gene causes elevated IOP and hypertensive glaucoma-like phenotypes, including degeneration of the optic nerve and RGCs and dysregulated visual functions. Regarding the mechanism involved in IOP reduction, the P2Y₆ receptor in the NPE cells suppressed the production of aqueous humor from the CB.
Figure 5. Reduction of IOP prevents RGC damage. (A) Daily topical administration of latanoprost to P2Y6KO mice over 4 months reduced the basal IOP to that of WT mice (n = 10–20, *P < 0.05 vs. WT 6 months, **P < 0.01 vs. P2Y6KO 6 months, 1-way ANOVA followed by Fisher’s LSD test). (B) Thinning of ganglion cell layer and inner plexiform layer (GCL/IPL) in 6-month-old P2Y6KO mice observed by OCT was restored by daily latanoprost administration (n = 10–20, *P < 0.05, **P < 0.01 vs. WT 6 months, ***P < 0.01 vs. P2Y6KO 6 months, 1-way ANOVA followed by Fisher’s LSD test). (C and D) Latanoprost prevented the reduction in RGC number in 6-month-old P2Y6KO mice (n = 25, **P < 0.01 vs. WT 6 months, ***P < 0.01 vs. P2Y6KO 6 months, 1-way ANOVA followed by Fisher’s LSD test). Values are the mean ± SEM. Scale bars: 100 μm in B and C, 250 μm in B.
Nucleotides have been reported to be present in the aqueous humor (33) and released from various eye tissues and cells (9, 34–37) in response to physiological stimuli; for example, physiological IOP elevation evokes exocytotic ATP release from the CB (12, 38). Pintor’s group has extensively studied nucleotide-mediated IOP changes (18–20, 39), and our data showing a hypotensive UDP effect was consistent with their results. In addition to the physiological roles of extracellular nucleotides, intraocular nucleotide levels have been reported to be elevated in the aqueous humor of glaucoma patients (13, 14, 40), indicating the pathological roles of dysregulated purinergic signaling in glaucoma. However, before this study, it was unclear whether the dysregulation of purinergic signaling contributed to the pathogenesis of glaucoma.

Histological analysis revealed that P2Y$_6$ receptors are localized in NPE cells and stroma of the ciliary processes, similar to a previous report (16). Indeed, Ca$^{2+}$ imaging showed that primary cultured NPE cells, but not PE cells, clearly responded to UDP. Because the CB mediates aqueous humor production, we used fluorophotometry with a minor modification to investigate the role of the P2Y$_6$ receptor. We used 2 conditions including Fph-A and -B. The Fph-B model showed a linear fluorescence increase in the anterior chamber, which was suppressed by UDP or timolol but not by latanoprost. P2Y$_6$KO mice showed an accelerated fluorescence increase, which was not inhibited by UDP. In contrast, the Fph-B model showed a fluorescent reduction, which was enhanced by latanoprost but not by UDP, timolol, or P2Y$_6$KO. Thus, we concluded that P2Y$_6$ receptor controls the dynamics of the aqueous humor in a similar way to timolol. Because timolol suppresses the production of the aqueous humor (41–44), the mechanism involving P2Y$_6$ receptors might be mediated by a similar mechanism. If this is true, how does the P2Y$_6$ receptor control the production of the aqueous humor? Transepithelial Cl$^-$ transport across the ciliary epithelium was reported to produce the driving force for aqueous humor formation (45–50). Cl$^-$ secretion from NPE cells is mediated by Cl$^-$ channels, whose activity is controlled by intracellular cyclic adenosine monophosphate (cAMP), which is increased by P2Y$_6$ receptor activation (51, 52). It has been reported that cAMP causes both an increase (52–54) and decrease of Cl$^-$ secretion (55–57). One explanation for these controversial reports might be related to TGFβ-mediated mechanisms. TGFβ reverses cAMP-mediated Cl$^-$ channel activation (58). Because P2Y$_6$ receptor activation causes TGFβ production (59, 60), UDP may suppress Cl$^-$ secretion by producing both TGFβ and cAMP. Another possible candidate is the gap junction because it mediates intercellular connections between PE and NPE cells, which is essential for transepithelial Cl$^-$ transport (56, 61). Elevation of cAMP caused intercellular uncoupling between PE and NPE cells (56). Elevated [Ca$^{2+}$]$^{ii}$ also induced the uncoupling of gap junctions (62–64). P2Y$_6$ receptor activation may suppress transepithelial Cl$^-$ transport via the uncoupling of PE and NPE cells by the elevation of cAMP and [Ca$^{2+}$]$^{ii}$. In addition to NPE cell-mediated mechanisms, P2Y$_6$ receptors are localized in the stroma of the ciliary process (16). As discussed in a previous study (20), because P2Y$_6$ receptors in blood vessels cause vasoconstriction (65, 66), the hypotensive effect by P2Y$_6$ receptors may be in part induced by vasoconstriction. These reports suggest that P2Y$_6$ receptor may suppress the production of the aqueous humor via multiple mechanisms.

Accelerated formation of the aqueous humor in P2Y$_6$KO mice suggested that IOP levels should be altered in these mutant mice. Indeed, P2Y$_6$KO mice showed a sustained elevation of IOP. CCT measurement revealed that P2Y$_6$KO mice showed thicker CCT at middle age. Many reports have suggested that IOP is positively correlated to CCT (67–75). Our present data demonstrated that mature adult P2Y$_6$KO
mice had elevated IOP compared with WT mice without differences in CCT; thus, we concluded that the elevated IOP value in mature adult P2Y6KO mice is not caused by CCT changes but rather by changes in aqueous humor flow. In contrast, we found that the CCTs of WT and P2Y6KO mice were increased at middle age. Human data have shown that a 1-mmHg IOP increase requires a 10–100 μm increase in CCT. Our results show an increase of IOP with 2.8 (WT) and 3.4 mmHg (P2Y6KO) in 3- to 12-month-old mice. Therefore, if these IOP increments were all caused by CCT changes, the CCTs at 12 months of age should be larger with 28–280 (WT) and 34–340 mmHg (P2Y6KO). Our CCT data showed increases of 6.5 (WT) and 29.4 μm (P2Y6KO); therefore, we concluded that the age-associated IOP increase was not caused by CCT changes but mainly by the altered dynamics of the aqueous humor.

We found that P2Y6KO mice showed structural and functional abnormalities of the retina at > 6 months old. Structural damage was reduced by lowering IOP with the daily application of latanoprost, indicating that elevated IOP is a cause of glaucoma-like phenotypes in P2Y6KO mice. We also report controversial results. First, WT mice at 12–18 months of age showed elevated IOP equivalent to that of P2Y6KO mice at 6 months. Despite the high IOP, WT mice showed no significant abnormalities. Second, P2Y6KO mice at 3 months of age already showed high IOP but no structural or functional abnormalities. These results suggest that IOP is one of a number of causes of glaucoma. What is the difference between WT and P2Y6KO mice? As we previously reported, the P2Y6 receptor is expressed in NPE cells and other cell types, including RGCs (76) and microglia (77). P2Y6 receptor in RGCs tonically enhances axonal function, and the KO of P2ry6 resulted in impaired axonal elongation (76). Because impaired axonal function is the first histological change in glaucoma (78), P2Y6 KO RGCs may have vulnerable axons and glaucoma-like phenotypes. P2Y6 receptors are essential for controlling microglial functions, and they were activated in an animal glaucoma model (79) and human glaucoma patients (80). Minocycline, an inhibitor of microglia, improved the integrity of the optic nerve head in a mouse model of glaucoma without a reduction in IOP (81). These reports indicate that aberrantly activated microglia might cause RGC damage as a downstream event of P2Y6 receptor deficiency.

We also found that P2Y6 receptors were downregulated in the CB, even in middle-aged WT mice. Possible candidates for this mechanism are PKCα (82) and prostaglandin E2 (PGE2) (83). Because these molecules increase with aging (84, 85), age-associated expression changes of P2Y6 receptor may be controlled by several signaling pathways. Associated with the expression level, the efficacy of UDP on IOP was also reduced in middle-aged WT mice. Thus, our data suggest that P2Y6 receptor function is disordered in an age-associated manner, thereby elevating IOP. Furthermore, the dysregulation of P2Y6 receptor function may increase the risk for the pathogenesis of glaucoma.

Taken together, our data demonstrate that P2Y6 receptor activation changes aqueous humor dynamics, thereby reducing IOP levels. The dysregulation of P2Y6 receptors causes sustained IOP elevation, aging-dependent RGC damage, and impairment in visual functions, which are similar to the phenotypes of hypertensive glaucoma. Therefore, the P2Y6 receptor is a key target for the treatment of glaucoma.

**Methods**

*Animals.* Mice were obtained from Japan SLC. P2Y6 receptor KO mice were generated as previously reported (86). Mice were kept at 23°C with a 12-hour (6:00 am to 6:00 pm) light-dark cycle. Experiments for IOP measurement and aqueous humor dynamics were performed between 6:00 pm and 12:00 pm.

*Chemicals and antibodies.* Reagents were obtained from the following sources. BSA, N,N’-1,4-Butanediylbis[N-(3-isothiocyanatophenyl) thiourea (MRS2578), paraformaldehyde (PFA), Triton X-100, and UDP were purchased from Sigma-Aldrich. DAPI was obtained from Dojindo. Sodium fluorescein (1.03887.0050) was purchased from Millipore. An anti-BRN3A antibody (goat polyclonal, sc-31984) was purchased from Santa Cruz Biotechnology. P2Y6 antibody (rabbit polyclonal, ab13391) was obtained from Abcam. Eye drops containing tropicamide (0.5%) with phenylephrine (0.5%) (Mydrin-P), latanoprost (0.005%, Senju), or timolol (0.5%, Teika) were purchased from Santen Pharmaceutical Co., Senju Pharmaceutical, and Teika Pharmaceutical, respectively.

*Measurement of IOP.* IOP was measured using a rebound-type tonometer (TonoLab). Mice were anesthetized with pentobarbital (i.p., 50 mg/kg), and IOP was measured after loss of righting reflex. IOP measurement was performed following the manufacturer’s protocol. The tonometer was placed 5 mm away from the eye at an angle of 90°. IOP was measured 5 times per day, and the values were averaged. All IOP measurements were performed from 6 pm to 12 pm. For eye drop treatment, 5 μl of agents were topically applied 30 minutes before IOP measurement.
Aqueous humor dynamics assays. Aqueous humor dynamics were measured by fluorophotometry. Mice were anesthetized with pentobarbital (50 mg/kg), and the dynamics were measured using 2 models including model A (Fph-A) and B (Fph-B). In Fph-A, sodium fluorescein (0.2%, 100 μl) was i.p. injected into mice, and fluorescence levels in the anterior chamber were monitored by a cooled CCD camera (3.2 megapixel Fujifilm super CCD) equipped with LAS-4000 (GE Healthcare) every 2 minutes. Because the IOP-lowering agents showed their maximum effect within a few hours (88), we set a test time of 30 minutes. In Fph-B, fluorescein (0.02%, 5 μl) was topically administered for 5 minutes, followed by washing 5 times with 500 μl of saline (89). UDP (500 μM, 5 μl), latanoprost (0.005%, 5 μl), timolol (0.5%, 5 μl) were applied in eye drops 30 minutes before fluorescein administration. Mice were anesthetized with sodium pentobarbital (50 mg/kg, i.p.) 25 minutes after eye-drop treatment. In Fph-A, the endogenous (without fluorescein injection) fluorescence level in the anterior chamber was equivalent to the level in the first images immediately after fluorescein injection (data not shown). The fluorescence level steadily increased to 300% at 30 minutes. In Fph-B, the fluorescence level was slightly increased during the first 4–6 minutes and then decreased, consistent with a previous report (89).

Preparation of whole-mount retinas. Eyes were enucleated and immersed in 4% of PFA for 30 minutes at room temperature and then dissected. Retinas were postfixed for 12 hours at 4°C. Immunostained retinas were mounted with Slow Fade Gold antifade reagent (Life Technologies).

Preparation of retinal sections. Enucleated eyes were fixed in 4% of PFA for 1 day at 4°C and soaked in 20% and 30% sucrose/PBS for 3 hours each. The samples were then embedded in Tissue-Tek optimal cutting temperature compound (Sakura Fintechanical) and frozen. All specimens were sectioned at 20 μm in the plane of the optic nerve head. For H&E staining, tissues were embedded in paraffin.

OCT. OCT was performed using a Micron IV Retinal Imaging Microscope (Phoenix Research Labs) following the manufacturer’s instructions. Mice were anesthetized with ketamine (100 mg/kg) and xylazine (9 mg/kg). Cross-sectional images of retinas were obtained from the region surrounding the optic nerve head (shown as green circles in the fundus images of Figure 4D and Figure 5B).

Real time RT-PCR. Total RNA was isolated and purified from astrocytes and neurons using RNeasy (Qiagen) according to the manufacturer’s instructions. Reverse transcription PCR (RT-PCR) was performed using a PrimeScript One Step RT-PCR Kit (Takara Bio Inc.) according to the manufacturer’s protocol. The reaction mix contained 40 ng total RNA, 200 nM primers, 100 nM TaqMan probe, TAKARA EX Taq HS, and PrimeScript RT enzyme Mix. RT-PCR amplification and real-time detection were performed using an Applied Biosystems 7500 Real-Time PCR System (Applied Biosystems). The reverse transcription was performed at 42°C for 5 minutes, followed by inactivation at 95°C for 10 seconds. The temperature profile consisted of 40 cycles of denaturation at 95°C for 5 seconds and annealing/extension at 60°C for 34 seconds. Primers and the Taqman probes for mouse Gapdh (catalog 4308313) and P2ry6 (Mm02620937_s1) were obtained from Applied Biosystems.

In situ hybridization. For in situ hybridization, 8-week-old male ICR mice were used. A 581-bp DNA fragment corresponding to nucleotide positions 11–591 of mouse P2ry6 (GenBank accession number NM_183168.2) was subcloned into pGEMT-Easy (Promega) and used to generate sense or antisense RNA probes. Sections were deparaffinized with xylene and rehydrated through an ethanol series and PBS. Sections were fixed in 4% PFA for 15 minutes and then washed with PBS. The sections were treated with 8 μg/ml proteinase K in PBS for 30 minutes at 37°C, washed with PBS, fixed with 4% PFA, washed again with PBS, and placed in 0.2 N HCl for 10 minutes. After washing with PBS, the sections were acetylated by incubation in 0.1 M triethanolamine-HCl (158917, MilliporeSigma) (pH 8.0, containing 0.25% acetic anhydride) for 10 minutes. After washing with PBS, the sections were dehydrated through an ethanol series. Hybridization was performed with probes at concentrations of 300 ng/ml for 16 hours at 60°C. After hybridization, sections were washed in 5× HybriWash (SHW-01, Genostaff, equivalent to 5× SSC) at 60°C for 20 minutes, then in 50% formamide, 2× HybriWash at 60°C for 20 minutes, followed by RNase treatment in 50 μg/ml RNase A in 10 mM Tris-HCl (pH 8.0, containing 1 M NaCl and 1 mM EDTA) for 30 minutes at 37°C. Then, the sections were washed twice with 2× HybriWash at 60°C for 20 minutes, twice with 0.2× HybriWash at 60°C for 20 minutes, and once with TBS/T (0.1% Tween 20 in TBS). After treatment with G-block (GB-01, Genostaff) for 30 minutes, the sections were incubated with anti-digoxigenin antibody (11093274910, Roche Diagnostics) conjugated with alkaline phosphatase (1:1,000) for 2 hours at room temperature. The sections were washed twice with TBS/T and then incubated in 100 mM NaCl, 50 mM MgCl₂, 0.1% Tween 20, and 100 mM Tris-HCl (pH 9.5). Development reactions were performed in NBT/BCIP solution (11681451001,
Roche Diagnostics) overnight, followed by washing with PBS. The sections were counterstained with Kernechtrot stain solution (N3020, MilliporeSigma), dehydrated, and then mounted with Malinol (20091, Muto Pure Chemicals Co. Ltd.).

**Immunofluorescence staining.** Samples were blocked with 2% goat serum in PBS containing 2% Triton X-100 (PBST) (064K0164, Roche Diagnostics) for 1 hour at room temperature. Then, samples were incubated with primary antibodies for 3 days at 4°C with gentle agitation. The sections were washed 3 times with PBST at room temperature for 10 minutes and incubated with secondary antibodies for 1 hour at room temperature. Alexa 488– or Alexa 546–conjugated secondary antibodies (1:1,000, A11034 and A11010, Molecular Probes) were used. Primary and secondary antibodies were diluted in blocking buffer and PBST containing 0.3% goat serum, respectively. Nuclei were stained with DAPI (100 μg/ml, Dojindo). Fluorescent images were acquired with a FV1200 laser scanning confocal microscope (Olympus) or BV-8000 (Keyence).

**mfERG.** mfERGs were recorded using a VERIS 6.0 system (Electro-Diagnostic Imaging) as previously reported (90). Mice were anesthetized with a mixture of xylazine (10 mg/kg) and ketamine (25 mg/kg; i.p.). Four- or 13-month-old mice were considered young or aged mice, respectively. The pupils were dilated with 0.5% phenylephrine hydrochloride and 0.5% tropicamide. The visual stimulus consisted of 7 hexagonal areas scaled with eccentricity. The stimulus array was displayed on a high-resolution black and white monitor driven at a frame rate of 100 Hz. The second-order kernel, impaired in glaucoma patients (91), was analyzed.

**SBF-SEM.** Enucleated eyes were immersed in 0.1 M PBS containing 4% PFA (163-25983, Wako) and 0.5% glutaraldehyde (077-06271, Wako) for 3 hours at room temperature. Then, the hemisphere including the cornea and CB were dissected and further fixed for 12 hours at 4°C. Optic nerve sections were cut on a vibratome (Leica), and the specimens were stained with 0.4% OsO4, uranyl acetate, and lead aspartate, followed by embedding in Epon resin (Electronic Microscopy Sciences). SBF-SEM images were obtained using a Sigma VP SEM (Carl Zeiss).

**Analysis of cornea structures.** Enucleated eyes were fixed with 4% PFA for 1 hour at room temperature, and corneas were dissected. Specimens were stained with DAPI (10 μg/ml in PBS) for 1 hour at room temperature and mounted on a glass slide. Cell layers were imaged by z-stack imaging using an FV-1200 laser scanning confocal microscope (Olympus).

**Primary cultured epithelial cells from the CB.** Enucleated eyes were immediately immersed in PBS, and the ciliary bodies were isolated. The tissue was then sliced using a McIlwain tissue chopper (Mickle Laboratory Engineering). The slices were attached to a μ-dish (ibidi GmbH) and cultured in DMEM (11885-084, Gibco) supplemented with 10% FBS, 100 units/ml of penicillin, and 100 μg/ml of streptomycin under 10% CO2 at 37°C with medium changes every few days. NPE cells were identified as transparent cells under visible light images. PE cells contained brown depositions.

**Ca2+ imaging.** Ca2+ imaging was performed as previously reported with minor modifications (76, 92). Briefly, culture medium was replaced with balanced salt solution (BSS) of the following composition (in mM): NaCl 150, KCl 5.0, CaCl2 1.8, MgCl2 1.2, HEPES 25, and D-glucose 10 (pH 7.4). Cells were loaded Briefly, culture medium was replaced with balanced salt solution (BSS) of the following composition (in mM): NaCl 150, KCl 5.0, CaCl2 1.8, MgCl2 1.2, HEPES 25, and D-glucose 10 (pH 7.4). Cells were loaded
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