Intraocular pressure elevation is a major causal risk factor for glaucoma. The resistance to aqueous humor (AQU) outflow is a key determinant of IOP. The major site(s) of resistance to AQU drainage have not been determined precisely but are thought to be within or close to the inner wall of the Schlemm’s canal (SC). Along with the trabecular meshwork (TM), Schlemm’s canal is a critical structure in the pressure-dependent AQU drainage pathway. Increased resistance to drainage and pathologic IOP elevation are thought to result from direct or indirect insults that cause dysfunction of SC/TM cells. Much remains to be learned about the genes and molecular pathways that contribute to normal outflow, and to the etiology of increased outflow resistance underlying high IOP. The molecular mechanisms mediating outflow and the precise route(s) of AQU flow across the SC inner wall need further clarification. Paracellular and transcellular routes of AQU outflow have been proposed based on observation of pores, between or within cells, in fixed or frozen tissue. The route of flow has not been visualized in real-time in living tissue.

Mouse outflow physiology is remarkably similar to that of humans in terms of conventional (pressure-dependent) outflow with similarities in IOP values, SC and TM anatomy, and lack of “washout” (defined as the increase in outflow that occurs during prolonged ocular perfusion as happens in other nonhuman species). This makes the mouse a powerful model for determining the mechanistic basis of outflow. The mouse’s relatively small size combined with the wealth of genetic and physiologic tools available, and the relatively low costs of maintenance make the mouse an ideal system for large-scale screens for drugs affecting outflow.

A key parameter in the study of outflow physiology is the outflow facility (C). Currently, mouse facility is measured by cannulation either in vivo or in enucleated eyes. Measurement of C by cannulation is subject to several physical factors (hydration and temperature) that must be carefully monitored. In addition, microleaks around the needle at the point of entry are a concern. In live mice, eyeball movements and twitching can cause the needle to come out of the eye, resulting in failed experiments. The bulky nature of the experimental setup precludes studying pressure-dependent flow in real time by live imaging under high-resolution microscopes. There is a need for a versatile system where...
We report the development of a perfusion system for mouse culture, and live imaging in a powerful experimental animal. One can measure \( C \), culture eyes, and perform live imaging of the intact outflow pathway.

One attractive solution is the use of an in vitro outflow perfusion apparatus. This type of apparatus has been used to measure outflow in eyes of humans and other species, and to culture eyes, and perform live imaging of the intact outflow pathway. One can measure \( C \), culture eyes, and perform live imaging of the intact outflow pathway.

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**Materials and Methods**

**Animals**

All experiments were performed in compliance with the Association for Research in Vision and Ophthalmology (ARVO) Statement for the Use of Animals in Ophthalmic and Vision Research and the approval of The Jackson Laboratory Institutional Animal Care and Use Committee (IACUC). All mice (3–5 months old and matched sexes) were obtained through The Jackson Laboratory. The mice were kept on a standard 14-hour light/10-hour dark cycle with food and water available ad libitum. The mouse strains used are shown in the Supplementary Methods.

**Perfusion Apparatus**

A mouse eye perfusion apparatus was engineered based on a human anterior segment perfusion apparatus (Fig. 1, Supplementary Fig. S1). Our apparatus, milled from Lexan, consisted of a base with a conical frustum for mounting eyes (Fig. 1, Supplementary Fig. S1). The top of the frustum had a diameter of 2.7 mm and the base 3.4 mm. We built our first system based on the human apparatus, which had a large footprint (Supplementary Fig. S1). We then redesigned that base to reduce the footprint and for ease of manufacture (Fig. 1). Most importantly the frustum design and performance were identical in both devices (Supplementary Fig. S2).

**Mounting a Dissected Eye on Perfusion Apparatus**

A dissected eye (see Supplementary Methods) was kept moist, and mounted carefully and gently onto the frustum using the opening in the side of the globe. The flaps cut into the sclera allowed the eye to sit on the frustum, and the flaps were lightly glued onto the platform bearing the frustum (using tissue glue) to hold the eye in position. A seal was created between the eye and frustum using an O-ring and washer (Fig. 1). The washer was secured in place by four corner screws that insert into the base. For uniformity, all screws were symmetrically and equally tightened using a torque screwdriver (10 cN.m; Fig. 1).

In this configuration the limbus of the eye spans the top of the frustum with sclera on one side and cornea on the other side. Although the retina is removed, the retinal pigment epithelium layer is maintained intact and lines the sclera. The eye was checked for a pressure-tight seal by gently pushing in test perfusate (Dulbecco's PBS [DPBS], balanced salt solution [BSS], or a drug mixture) through the inlet port, after closing the outlet port, using a three-way stopcock to determine if the flaccid eye mounted on the frustum inflated without visible leaks around the O-ring.

The inlet port then was connected to a pressure transducer (DTXPlus TNF-R; Argon Medical Systems, Plano, TX, USA; see Supplementary Methods for details), in series with a perfusate-filled 100 μl Nanofil syringe (Worcester Polytechnic Institute [WPI], Worcester, MA, USA) attached to a UMP3 pump (WPI) controlled by a Micro-i pump controller (WPI). The transducer was connected to the apparatus with a short length of pressure-resistant tubing. Any air bubbles in the system were removed carefully using the perfusion fluid of choice. The pressure transducer connects to the computer via a digitizer (Measurement Computing, Norton, MA, USA) and with the microprocessor controller we created closed-loop control (WPI; see below). The eye was maintained in a moist state at 37°C by inverting the eye over a well of a heating block filled with water (to a height approximately 1 mm below the eye, see Fig. 2B) during measurement of outflow.

**FIGURE 1.** Schematic of the perfusion apparatus with a mounted eye. (A) The device consists of a base with central frustum onto which the eye is mounted. The frustum has two fluid ports (see B) for addition and removal of fluids. The eye is held in place using an O-ring that seals the eye against the frustum. A mounting washer and 4 screws hold the O-ring in place creating a tight seal. (B) A cross-sectional view of the system. The mounted eye is oriented so that approximately 45% of the limbal tissue, including the drainage structures, is clear of where the O-ring clamps the tissue. Approximate position of the optic nerve (ON) stub is indicated. Images not to scale.
Monitoring and Control System

While measuring outflow, fluid was perfused at a controlled rate to maintain the eye at a specific pressure. The perfusion rate was monitored and controlled by an in-house computer program (LabVIEW; National Instruments, Austin, TX, USA) using a bang-bang control algorithm. The steady state error of the system was ±0.5 mm Hg. Once started, the program continuously monitored and recorded information from the pressure transducer and flow rate information from the syringe pump microprocessor controller. If pressure fell by a small fraction of 1 mm Hg below the set level, the program triggered the syringe pump to perfuse fluid into the eye via the microprocessor controller. As soon as the pressure increased to a fraction of 1 mm Hg above the set point the pump turned off. As the pump delivered fluid in increments of 0.67 nL/step, flow continued after the pump was shut off, explaining the oscillatory nature of the flow rate plot. The software was designed to set different experimental pressure levels in a stepwise manner (up to six pressure values can be input). The user set the “rise” time to reach a set pressure, and a “record” time to measure flow rate required to maintain the pressure. The rise and equilibration time ensured that we recorded data only after the flow rate oscillations had reached a stable state.

Outflow Measurements

Before outflow measurement on any day, we calibrated the pressure transducer using a mercury manometer. Outflow was measured by obtaining the flow rate required to maintain a series of set pressures. We use two pressure series: Protocol A – 5, 15, 25, 35, 45, and then 15 mm Hg again, and Protocol B – 8, 12, 16, 20, 24, and 12 mm Hg again. In both protocols, rise time was 5 minutes, equilibration time was 5 minutes, and record time was 10 minutes (see Monitoring and control section above). A run was considered successful if the following criteria were met in analysis of the data: (1) Flow rates at each pressure achieved stable oscillation about a mean with oscillations being no more than ±30 nL for 5, 15, and 25 mm Hg pressure set points, and (2) Flow rates were similar at the...
first and second 15 mm Hg (Protocol A) or first and second 12 mm Hg (Protocol B). Our success rate was over 90%. If an eye failed at any pressure, all data for that eye were discarded. Drugs were perfused in at 15 mm Hg for 30 minutes followed by 15 minutes at 5 mm Hg before measuring outflow. We used Y27632 at 200 μM and L-NAME at 100 μM. No eyes were perfused for longer than 2 hours and some eyes were fixed for immunofluorescence or histology (see Supplementary Methods) after outflow measurements were concluded.

RESULTS

The Perfusion System

The system consisted of a perfusion platform with a conical frustum for mounting the eye (Fig. 1, Supplementary Fig. S1) and a micropump fluid perfusion system controlled by a computer (Fig. 2A). We first determined the best way to mount eyes onto the frustum. Due to mouse anatomy, mounting the anterior segment of the eye “cornea up,” as with human anterior segments, placed the limbus too close to all clamping mechanisms that we devised. This risked crushing the drainage structures and blocking outflow. Thus, we mounted the eye with the limbus up and spanning the top of the frustum (therefore, ~45% of the limbal circumference was available for perfusion; see Supplementary Methods). This approach provided ample clearance from the O-ring and an orientation that would ultimately facilitate live imaging. To enable this, we developed the dissection scheme that is shown in Supplementary Figures S3 and S4 (see Supplementary Methods). The dissection of the eye is a critical and delicate step that requires attention to detail.

Characterization of the System: System Compliance and Resistance

Before performing any outflow measurements, we tested the system for any inherent resistance and compliance (defined as change in flow rate in response to pressure caused by any elasticity in the system). It is critical to determine these parameters because they can make outflow measurement inaccurate. Upon using a flow rate of 660 μl/min (far larger than the flow observed by an eye) and leaving the inlet port open (no eye secured to system), we observed no increase over atmospheric pressure (0 mm Hg), indicating that the system had no measurable resistance at this flow rate (or under physiologic conditions). System compliance was determined to be 20.65 nL/mm Hg (see Supplementary Methods).

Washout in Mouse Eye

Washout (C increase upon prolonged ocular perfusion) is observed in some nonhuman species, including primates but not humans or mice.10,22–25 The washout rate for mouse eyes using our system was 2.5 ± 1.96% (n = 6), in agreement with an earlier study (washout rate = 2.4%)10 and in contrast to monkey eyes, where the washout rate is 26% to 42%.24 Thus, we confirmed that mouse eyes, like human eyes,25 have negligible washout.

Outflow Measurements

We determined C in mouse eyes by computing the slope of the flow rate versus pressure plot (Protocols A and B, see Supplementary Methods).10 An example of recorded pressure and flow rate traces is shown in Figure 3. Importantly, this plot shows that the flow rates at 15 mm Hg at the beginning and end of the perfusion protocol are similar, indicating that the
In Vitro Perfusion System for Mouse Eye Outflow Studies

Inhibition of Rho-Associated Protein Kinase (ROCK) Increases Outflow

We next determined if perturbation of outflow at a molecular level can be detected using our perfusion device. We first tested the effect of the ROCK inhibitor, Y-27632 (see Supplementary Methods for details), on C. Y-27632 is well established to increase outflow in other species. We observed a 74.4% increase in C in eyes treated with Y-27632 (n = 10, t-test P = 0.003; Fig. 6). Thus, as expected, ROCK inhibition increased C. This experiment validated the use of our system to measure changes in C.

Nos3 and Cav1 Are Required for Proper Outflow

Nos3 (endothelial nitric oxide synthase, eNOS) and Cav1 are genes implicated in glaucoma. It is unclear how these genes impact the disease. Hence, we determined if these genes were required for normal outflow. We first disrupted Nos3 function by inhibition using the inhibitor L-NAME; see Supplementary Methods for details) as well as through genetic mutation. Treatment with L-NAME caused a reduction in C by 28.3% from control (n = 7 eyes; t-test, P = 0.025; Fig. 7A). In Nos3 null mutant eyes C was reduced by 35.8% compared to control (n = 7 eyes; t-test, P = 0.008; Fig. 7B). Next, we performed outflow measurements on Cav1 null (Cav1−/−) eyes (n = 9 eyes; Fig. 8). In Cav1−/− eyes, C was reduced by 47.8% compared to control (t-test, P = 0.02). We detected this reduction in two independent experiments on a B6 strain background and on a mixed strain background (not shown). These results indicated that NO3S and CAV1 function are required for normal outflow.
DISCUSSION

We described a new way to study the molecular mechanisms controlling outflow. Using a newly engineered in vitro perfusion system, we first established a mechanistic role for the kinase activity of ROCK in mouse conventional outflow. We also showed that NOS3 and CAV1 function, both previously implicated in glaucoma, is necessary for normal outflow. These studies demonstrated the use of our system for identifying molecules and/or pathways controlling outflow.

Conventional facility measured using our system was in the range described for mouse eyes. Several other investigators have described the nonlinearity of the flow rate–pressure plot above 25 mm Hg. Recently, it has been determined that anterior chamber deepening with increasing pressure (over 25 mm Hg) in enucleated mouse eyes results in traction on the TM, thus increasing C. Our mounted eyes do not have an intact iris and AC, but similar nonlinearity was observed. Although the mechanism explaining this is not clear, increasing traction on the mounted tissue may itself allow nonlinear increases in outflow with increasing pressure. A recent publication, using closely spaced pressure points and sophisticated data analysis, supports nonlinear changes in outflow with changing pressure.

A recent report suggested that insufficient hydration results in a positive β-intercept and that hydrating enucleated eyes by immersion eliminated this positive β-intercept, actually rendering it negative. During our experiments, we had maintained...
the mounted eyes in a humid environment (measured to be 100% humidity) to prevent evaporation. Nevertheless, our positive intercepts suggested that the humidity measurements might not have been accurate, allowing some evaporation. Thus, in response to this recent report, we performed our perfusion protocol after submerging eyes in PBS at 37°C. The submersion changed the y-intercept to ~18.5 μL/min, indicating that evaporation from the surface of the eye may be occurring in our humid environment. Importantly, the C value obtained when submerging the eyes was $C = 5.065 \pm 1.1$ nL/mm/Hg. This value is not significantly different from that obtained when eyes were maintained in a humid atmosphere ($4.7 \pm 1.2$ nL/mm/Hg; $P = 0.49$). This suggested that our original approach of maintaining the eye in the humid atmosphere did not affect measurement of C. Importantly and during peer review of the current study, another study found no flow at 0 mm Hg and suggested that negative intercepts are an artifact resulting from linear fitting of data.35 This study sampled flow at nine pressures over the 5 to 20 mm Hg range. This dense sampling allowed power-fitting with a y-intercept of zero at 0 mm Hg. Such dense sampling can be tested with our perfusion system to study molecular mechanisms controlling outflow.

Cav1 has been implicated in glaucoma.30 Given the ubiquitous expression of CAV1, it could be important for IOP, function of retinal ganglion cells, and cells in the optic nerve head. Our data indicated that Cav1 is at least required for conventional drainage and will impact IOP. Given that we used a mutant mouse where Cav1 is globally deleted, we will in the future perform cell type-specific inhibition to identify the critical cell type that controls outflow through Cav1 function.

The protein Cav1 has been implicated in mechanotransduction and regulation of Nos3 activity.36 Loss of either function (not mutually exclusive) would prevent translation of pressure-dependent strain on SC/TM cells into signals required for outflow.

In summary, we have successfully developed a new perfusion system to study molecular mechanisms controlling outflow. We described a new role for Cav1 in outflow physiology and provide more evidence for central roles for
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![Flow-rate versus pressure graph](image)

**FIGURE 8.** Loss of Cav1 decreases pressure-dependent outflow in mouse eyes. Flow-rate versus pressure graph showing a comparison of flow rates in control and Cav1<sup>−/−</sup> eyes. The equations representing the trend line for the linear part of the curve were: Control, \( y = 0.007x + 0.026 \), \( R^2 = 0.99322 \), black circles; Cav1<sup>−/−</sup>, \( y = 0.005x + 0.039 \), \( R^2 = 0.99322 \), black diamond. SEM is shown for data points in the graph. The computed \( C \) values were: control, \( C = 6.9 \pm 3.4 \text{nL/min/mm Hg} \) (whole eye \( C = 15.18 \pm 4.4 \text{nL/min/mm Hg} \), and \( C = 3.6 \pm 2.5 \text{nL/min/mm Hg} \) (whole eye \( C = 7.92 \pm 5.5 \text{nL/min/mm Hg} \)). \( C \) = mean \pm SD. SEM is shown for data points in the graph.

NOS3 and ROCK in outflow regulation. With its potential for organ culture, our system is expected to allow development of longitudinal physiologic studies and live imaging.

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