





Cell atlas of aqueous humor outflow pathways in eyes of humans and four model species provides insight into glaucoma pathogenesis

Tavé van Zyl^{a,b,c,1,2} , Wenjun Yan^{b,c,1}, Alexi McAdams^{a,b,c}, Yi-Rong Peng^{b,c,3}, Karthik Shekhar^{d,e,4}, Aviv Regev^{d,e,f}, Dejan Juric^g, and Joshua R. Sanes^{b,c,2} 

^aDepartment of Ophthalmology, Harvard Medical School and Massachusetts Eye and Ear, Boston, MA 02114; ^bCenter for Brain Science, Harvard University, Cambridge, MA 02138; ^cDepartment of Molecular and Cellular Biology, Harvard University, Cambridge, MA 02138; ^dHoward Hughes Medical Institute, Massachusetts Institute of Technology, Cambridge, MA 02142; ^eKoch Institute of Integrative Cancer Research, Department of Biology, Massachusetts Institute of Technology, Cambridge, MA 02142; ^fKlarman Cell Observatory, Broad Institute of Massachusetts Institute of Technology and Harvard, Cambridge, MA 02142; and ^gDepartment of Medicine, Harvard Medical School and Massachusetts General Hospital Cancer Center, Boston, MA 02114

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Increased intraocular pressure (IOP) represents a major risk factor for glaucoma, a prevalent eye disease characterized by death of retinal ganglion cells; lowering IOP is the only proven treatment strategy to delay disease progression. The main determinant of IOP is the equilibrium between production and drainage of aqueous humor, with compromised drainage generally viewed as the primary contributor to dangerous IOP elevations. Drainage occurs through two pathways in the anterior segment of the eye called conventional and uveoscleral. To gain insights into the cell types that comprise these pathways, we used high-throughput single-cell RNA sequencing (scRNAseq). From ~24,000 single-cell transcriptomes, we identified 19 cell types with molecular markers for each and used histological methods to localize each type. We then performed similar analyses on four organisms used for experimental studies of IOP dynamics and glaucoma: cynomolgus macaque (*Macaca fascicularis*), rhesus macaque (*Macaca mulatta*), pig (*Sus scrofa*), and mouse (*Mus musculus*). Many human cell types had counterparts in these models, but differences in cell types and gene expression were evident. Finally, we identified the cell types that express genes implicated in glaucoma in all five species. Together, our results provide foundations for investigating the pathogenesis of glaucoma and for using model systems to assess mechanisms and potential interventions.

trabecular meshwork | ciliary muscle | macaque | pig | mouse

Glaucoma, a leading cause of irreversible blindness worldwide (1), results from loss of retinal ganglion cells (RGCs), which carry information about the visual world from the eye to the rest of the brain (2). Of the major risk factors, including age, race, and family history, the only modifiable one is intraocular pressure (IOP). Indeed, lowering IOP remains the only proven treatment strategy to delay disease progression, with several pharmacological and surgical approaches in widespread clinical use (2). However, therapeutic advances have been limited by incomplete understanding of the tissues that regulate IOP and the means by which increased IOP leads to RGC loss. We address the first of these issues here.

There are two major compartments in the eye, an anterior segment containing aqueous humor (AH) and a posterior segment containing vitreous humor. The primary determinant of IOP is the equilibrium between the production and drainage of AH. AH is produced by the ciliary body, circulates within the anterior chamber, and is then drained through one of two pathways, conventional (trabecular) or uveoscleral (3) (Fig. 1). In the conventional pathway, AH exits the anterior chamber through a lattice-like biological filter called the trabecular meshwork (TM), which is composed of collagenous beams lined with specialized TM cells. It then passes through a region of cells embedded within a denser extracellular matrix called the

juxtacanalicular tissue (JCT) prior to being conveyed into a specialized vessel called Schlemm canal (SC). From SC, AH exits the eye through a network of collector channels (CCs) continuous with the venous system. The remaining AH exits the anterior chamber via the uveoscleral pathway, draining through the interstices of the ciliary muscle (CM) and ultimately exiting the eye via the suprachoroidal space and the sclera. In principle, increased IOP could result from either excessive production or insufficient drainage of AH, but in practice, the latter is more commonly implicated (4–7).

The cellular composition of the AH outflow pathways has been studied histologically (8, 9), but because of their small size, intricate architecture, and tightly invested tissues, it is impractical to dissect individual components for molecular analysis.

Significance

Glaucoma is a leading cause of irreversible blindness worldwide. A major risk factor is increased intraocular pressure (IOP), and all current treatment strategies for slowing disease progression center on IOP reduction. Increased IOP generally results from reduced drainage of aqueous humor through two routes in the anterior segment of the eye called the conventional and uveoscleral outflow pathways. We use high-throughput single-cell RNA sequencing to identify cell types within these pathways and the genes that they express both in humans and in four model species frequently used to study IOP regulation and glaucoma.

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J.R.S. is a consultant at Biogen. All other authors declare no competing interest.

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Data deposition: The raw and unprocessed data files reported in this paper have been deposited in the Gene Expression Omnibus (accession no. [GSE146188](#)). Data can be visualized in the Broad Institute's Single Cell Portal at https://singlecell.broadinstitute.org/single_cell/study/SCP780.

¹T.v.Z. and W.Y. contributed equally to this work.

²To whom correspondence may be addressed. Email: tavevanzyl@gmail.com or sanesj@mcb.harvard.edu.

³Present address: Department of Ophthalmology and Jules Stein Eye Institute, University of California, Los Angeles, CA 90095.

⁴Present address: Department of Chemical Engineering, University of California, Berkeley, CA 94720.

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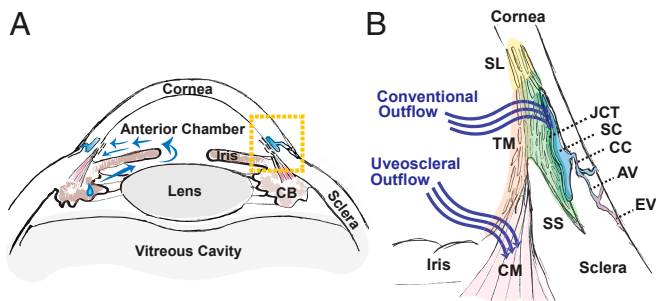


Fig. 1. Human AH outflow pathways. (A) Diagram of the anterior segment of the human eye, which includes the cornea, iris, ciliary body (CB), and lens. AH is secreted by the CB and circulates (blue arrows) within the anterior chamber prior to draining from the eye through one of two pathways located within the iridocorneal angle delineated by boxed area. (B) Enlarged view of the iridocorneal angle (boxed area in A) highlighting two outflow pathways for AH. In the conventional pathway, AH traverses the TM, first through the uveal meshwork (orange highlight), then the corneoscleral meshwork (light green highlight), and finally, the JCT (dark green highlight) prior to entering SC. AH exits the SC via CCs that empty into aqueous veins (AVs) that themselves merge with episcleral veins (EVs). Nonfiltering TM is located at the insert region (yellow highlight), referred to as Schwalbe line (SL), which abuts the corneal endothelium. In the uveoscleral pathway, AH exits via the interstices of the CM. SS, scleral spur.

Therefore, most studies of gene expression in these tissues have relied on bulk measurements of tissue or cultured cells (10–13). It has, therefore, been difficult to assess gene expression patterns of individual cell types. To circumvent this limitation, we used high-throughput single-cell RNA sequencing (scRNAseq) (SI Appendix, Fig. S1A). We profiled ~24,000 single cells from adult human TM and surrounding tissues, applied computational methods to cluster the cells based on their transcriptomes, and used histological methods to match the molecularly defined clusters to specific cell types. We identified 19 cell types and defined molecular markers for each of them.

We then used this cell atlas in two ways. First, we assessed expression of genes that have been implicated in glaucoma either as causal genes with Mendelian inheritance or as susceptibility loci identified in genome-wide association studies (GWAS) (14–22), and we compared expression levels in cell types of the outflow pathways with those in RGCs and retinal glia (23). We found that while many genes were preferentially expressed in the anterior segment, others were expressed predominantly in the retina.

Second, we used the human atlas as a foundation for assessing four animal models frequently used to study AH outflow pathways and glaucoma—cynomolgus macaque (*Macaca fascicularis*), rhesus macaque (*Macaca mulatta*), pig (*Sus scrofa*), and mouse (*Mus musculus*) (24–26). The utility of these models for disease research depends in large part on the extent to which their cell types and patterns of gene expression correspond to those in humans. We show that broad cell classes were generally conserved across all five species and could be localized to expected areas within the eye but that some cell types within classes were less well conserved. In some cases, molecular markers specific for human cell types were either absent or much less specific for cell types within other species. Disease genes in humans, which often demonstrated highly specific expression in outflow pathway cell types, mapped well in macaque species but less reliably in the pig and mouse. These results may help guide tests of therapeutic strategies in these models.

Results

Cell Atlas of Human Trabecular Meshwork and Aqueous Outflow Structures. We used a droplet-based method (27) to obtain 24,023 high-quality single-cell transcriptomes from human TM

tissue dissected from seven eyes of six postmortem donors (SI Appendix, Fig. S1B and Table S1). Computational analysis (SI Appendix) divided these cells into 19 clusters (Fig. 2A), ranging in frequency from 0.1 to 36% of cells profiled (Fig. 2C). Cells in nearly all clusters were obtained from all individuals (Fig. 2B) (the outlying clusters, C7 and C9, are discussed below). We identified differentially expressed genes for each cluster (Fig. 2D) and then used in situ hybridization and immunohistochemistry to relate the molecularly defined clusters to cell types defined by classical criteria of morphology and location.

Conventional pathway. The conventional pathway includes the TM, SC, CCs and closely associated structures (Fig. 1B). The TM includes an anterior, nonfiltering, insert region and a more posterior filtering region, which itself is divided into an inner part composed of “beam cells” through which AH first filters and an outer part composed of JCT cells residing adjacent to SC.

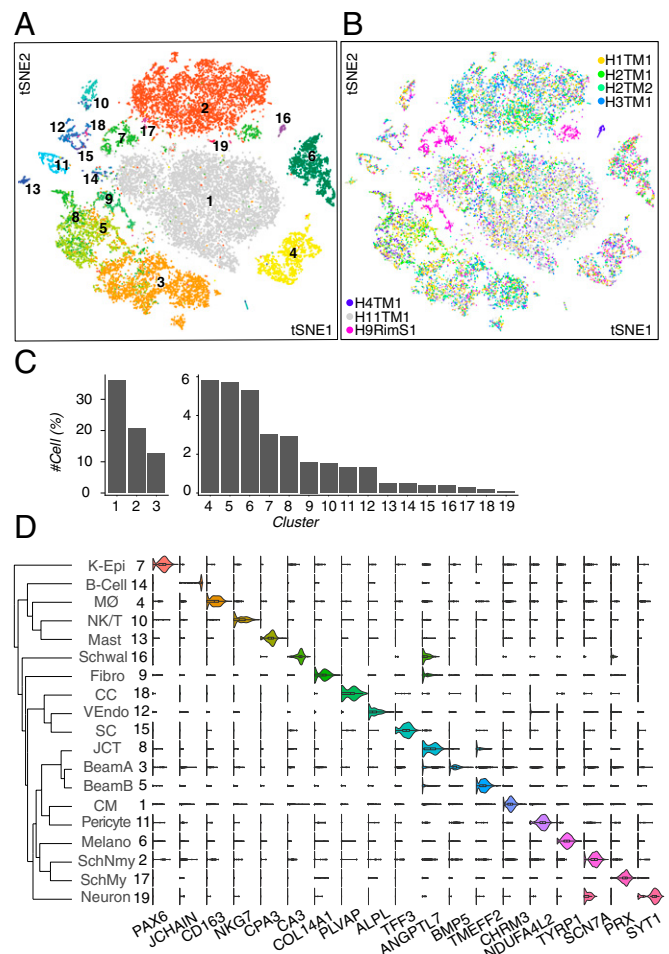


Fig. 2. Cell types and gene expression in the human outflow pathways. (A) Clustering of 24,023 single-cell expression profiles from human TM and associated structures visualized by t-distributed stochastic neighbor embedding (tSNE). Arbitrary colors are used to distinguish clusters deemed to be distinct by unsupervised analysis. Clusters were numbered according to relative size, with 1 being the largest. (B) tSNE plot shown in A but with cells colored by sample of origin. Note that corneal epithelial cells (C7) and fibroblasts (C9) were derived primarily from the rim sample (H9). (C) Frequency of each cell type; numbering is the same as in A and D. (D) Violin plots showing expression of genes selectively expressed by cells of each type. The dendrogram in Left shows transcriptional relationships among cell types. Fibro, fibroblast; K-Epi, corneal epithelium; Melano, melanocyte; MØ, macrophage; SchMy, myelinating Schwann cell; SchNmy, nonmyelinating Schwann cell; Schwal, Schwalbe line; VEndo, vascular endothelium.

These structures modulate outflow resistance through cell–cell interactions as well as through the generation and degradation of extracellular matrix (5, 7, 28–30).

We identified eight cell types within the conventional pathway, three of which corresponded to filtering TM cells. They were characterized and localized as follows.

Clusters 3, 5, and 8 demonstrated high expression levels of *MYOC*, *MGP*, and *PDPN* (Fig. 3A), markers previously associated with TM, as well as other genes, such as *RARRES1* (7, 31, 32). We used immunostaining for *PDPN* and *RARRES1* to confirm that these genes were expressed by cells encasing the trabecular beams (Fig. 3B and C and *SI Appendix, Fig. S2 K and L*). Clusters 3 and 5 could be distinguished by preferential expression of *FABP4* and *TMEFF2*, respectively, each of which marked a subset of beam cells, which we call Beam A and Beam B (Fig. 3D and *SI Appendix, Fig. S2 A, B, and J*). The *FABP4*+ Beam A cells and *TMEFF2*+ Beam B cells were intermingled but with a tendency for the latter to be closer to the JCT. Cluster 8 could be distinguished from C3 and C5 by selective expression of *CHI3L1* and *ANGPTL7*, both previously suggested to be potential markers for a TM cell subpopulation (13). In situ hybridization demonstrated localization of cells expressing these

genes predominantly to the JCT (Fig. 3E and *SI Appendix, Fig. S2I*). Other differentially expressed genes in this cluster (C8) included *RSPO4*, *FMOD*, and *NELL2*. Thus, our transcriptomic method identifies three cell types comprising the filtering TM: two types of beam cells and a distinct JCT cell.

A fourth cluster (C16) corresponded to cells of Schwalbe line, which is located in the “insert” region of the anterior nonfiltering meshwork abutting the peripheral corneal endothelium. This cluster had differentially expressed genes associated with corneal endothelium (e.g., *CA3*, *MGARP*, *SLC11A2*, *TGFBI*), but some cells shared markers associated with beam cells (e.g., *MYOC*, *IGFBP2*, *NELL2*, *PTGDS*). The candidate TM marker *AQP1* was preferentially expressed in these nonfiltering TM cells, consistent with the observation that deletion of *AQP1* in mice did not affect outflow facility (Fig. 3I) (33). Although it has been suggested that Schwalbe line contains TM precursors (34), we detected no differentially expressed genes suggestive of “stemness” (35) in this cluster; if present, they may have been too rare to be detected.

SC endothelial cells (C15) expressed canonical endothelial markers *PECAM-1* (*CD31*), vascular endothelial-cadherin (*CDH5*), and *CLDN5* as well as the hemostasis genes *PLAT* and *VWF* (Fig. 3A).

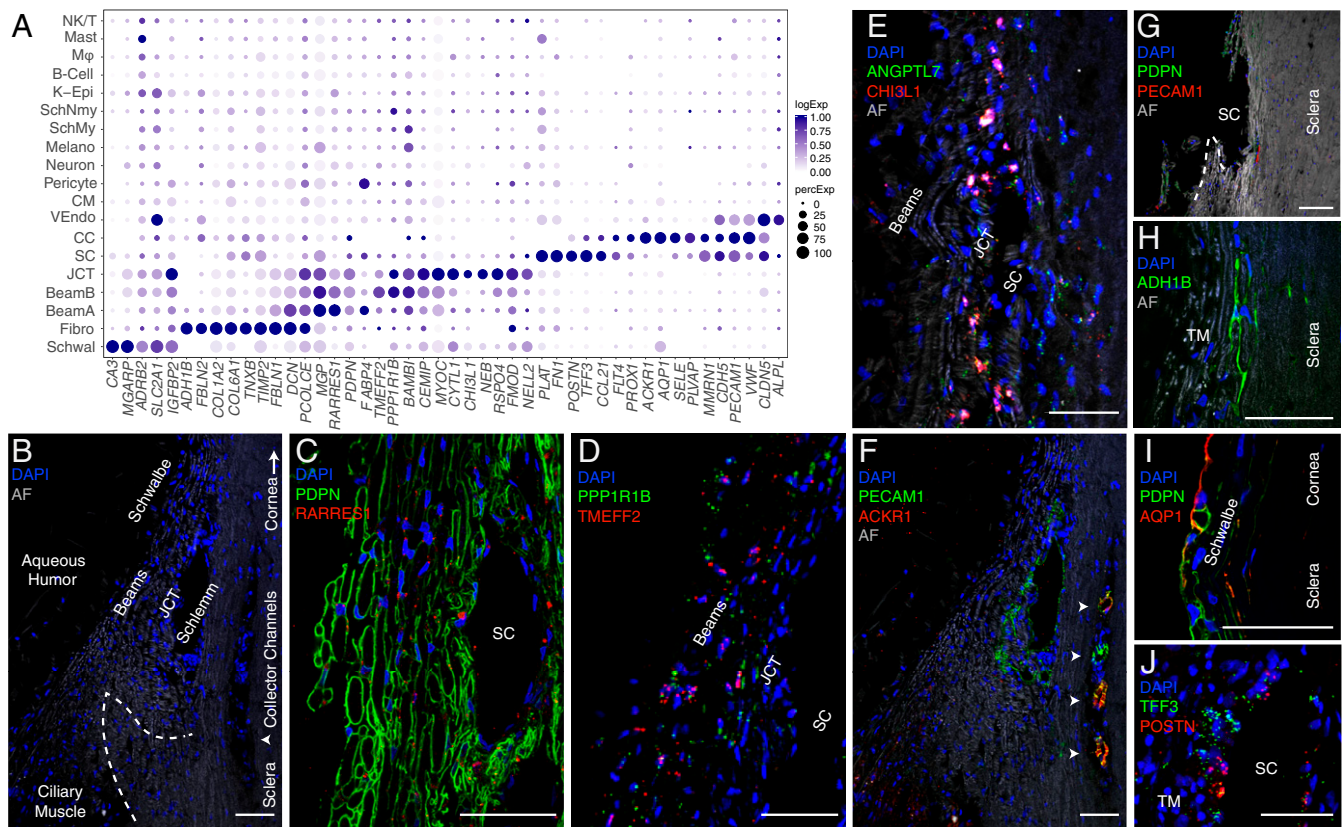


Fig. 3. Cells of the human conventional outflow pathway. (A) Dot plot showing genes selectively expressed in cells of the conventional outflow pathway. In this and subsequent figures, the size of each circle is proportional to the percentage of cells expressing the gene, and the shade intensity depicts the average normalized transcript count in expressing cells. Fibro, fibroblast; K-Epi, corneal epithelium; Melano, melanocyte; MØ, macrophage; SchMy, myelinating Schwann cell; SchNmY, nonmyelinating Schwann cell; Schwal, Schwalbe line; VEndo, vascular endothelium. (B) Labeled meridional section of a human corneoscleral rim visualized using autofluorescence (AF) demonstrates outflow anatomy at the iridocorneal angle visualized after removal of iris and ciliary body. The dashed line indicates scleral spur and arrowhead indicates region of CCs. (C) TM (beam and JCT) cells immunostained for PDPN (green) and RARRES1 (red). (D) Fluorescent RNA in situ hybridization against *TMEFF2* (red) and *PPP1R1B* (green) highlights beam cells. (E) Fluorescent RNA in situ hybridization against *ANGPTL7* (green) and *CHI3L1* (red) highlights cells in the JCT. (F) Immunostaining for PECAM1/CD31 (green) highlights SC, CCs (arrowheads), and CM capillaries, while ACKR1/DARC (red) contains only the CCs. The same field as in B. (G) Corneoscleral rim section demonstrates the tissue left behind after TM dissection. Very few cells in this region are positive for PDPN (green) or PECAM1/CD31 (red), indicating successful removal of relevant structures (TM and SC) during dissection protocol. (H) Scleral fibroblasts identified in the corneoscleral rim collection immunostained for ADH1B. (I) Schwalbe line cells at the junction of TM and corneal endothelium contain with PDPN (green) and AQP1 (red). (J) Fluorescent RNA in situ hybridization against *POSTN* (red) and *TFF3* (green) highlights SC endothelium. DAPI, 4',6-diamidino-2-phenylindole (nuclear stain). (Scale bars: 50 μ m.)

This cluster also selectively expressed the lymphatic endothelial cell markers *CCL21* and *FLT4* (*VEGFR3*), consistent with the notion that SC is a modified lymphatic vessel (36–38). Other lymphatic endothelial markers expressed in SC at lower levels were *PROX1* and *LYVE1*. Fibronectin-1 (*FNI*), a Transforming Growth Factor-beta-inducible extracellular matrix (ECM) gene implicated in glaucoma pathogenesis, was differentially expressed in SC endothelium as was *MMR1* (39, 40). We used the markers *POSTN* and *TFF3* (Fig. 3J) to distinguish endothelium of the outflow tract from the vascular endothelium in the CM, which shared many endothelial markers (discussed later). Some intermingled cells in SC expressed *PECAM-1* (*CD31*) and the vascular endothelial marker *ALPL* (Fig. 4C), raising the possibility that this structure contains two cell types.

Because blunt dissection of the TM may have left behind components of the conventional outflow structures such as the outer wall of SC, we collected a corneoscleral rim sample, which included all adjacent tissues after blunt dissection (Fig. 3G). This sample contained an additional vascular cell type (C18) and a fibroblastic type (C9) that were poorly represented in the other samples. Using the differentially expressed gene *ACKR1* (*DARC*), we identified C18 as the endothelium of the CCs downstream from SC (Fig. 3F). *ACKR1* has been identified as a marker to distinguish venular from nonvenular endothelial cells, which is consistent with the pre-/perivenular location of the channels (41). Other differentially expressed genes for CC cells include *SELE*, *SELP*, and *COL15A1*. The other cluster, C9, consisted of matrix fibroblasts. Immunostaining for the selectively expressed gene *ADHIB* showed that C9 cells were present within the sclera located adjacent to the TM and outer wall of SC (Fig. 3H and *SI Appendix*, Fig. S2H). This cluster was distinguished from TM cell types through its higher level of expression of *FBLN2*, *TIMP2*, *TNXB*, and multiple collagen genes (*COL1A2*, *COL6A1*, *COL6A2*, *COL6A3*, *COL14A1*), markers consistent with previous single-cell studies on fibroblasts in other organ systems (42). However, it also shared a set of ECM and complement genes with TM cells (C3, -5, -8), including *DCN*, *PCOLCE*, *FBLN1*, *MFAP4*, *SERPING1*, *CIS*, and *C1R*, suggesting that these neighboring cells play overlapping roles.

Finally, the corneoscleral rim sample included cells with transcriptomic signatures marking them as corneal epithelium (C7) based on expression of previously described markers, such as *AQP3*, *AQP5*, and *KRT5* (43). These cells will be described elsewhere.

Uveoscleral pathway. AH that does not exit the eye through the conventional pathway instead exits via the uveoscleral pathway, draining through the interstices of the CM. Seven clusters in our dataset comprised components of this pathway. The largest cluster (C1) corresponded to CM cells (44). It expressed markers classically associated with well-differentiated smooth muscle, including *DES*, *CNN1*, *MYH11*, *MYLK*, and *ACTC1* (Fig. 4A). We used the differentially expressed genes *DES* and *CHRM3* to localize this cluster histologically to the longitudinal CM (Fig. 4B and *SI Appendix*, Fig. S2C). The high expression levels of *ATP2A1/SERCA1*, a marker of type II fast twitch skeletal muscle, suggest that CM may also have characteristics of skeletal muscle, although it lacks expression of other classical skeletal muscle markers, including *MYH1*, -2, -4, and -7.

Six additional cell types were found within the CM. One, vascular endothelium (C12), was associated with intramuscular capillaries. This cluster shared canonical endothelial markers with SC and CCs but was distinguished from the latter by the presence of *ALPL* (Fig. 4C) and absence of lymphatic markers. Similarly, *PLVAP*, while present in SC and CCs, was not expressed in CM capillary endothelium, consistent with observations that these cells do not have pores (39, 45).

C11 comprises contractile pericytes, which express canonical markers *PDGFRB*, *MCAM/CD146*, and *NOTCH3* as well as

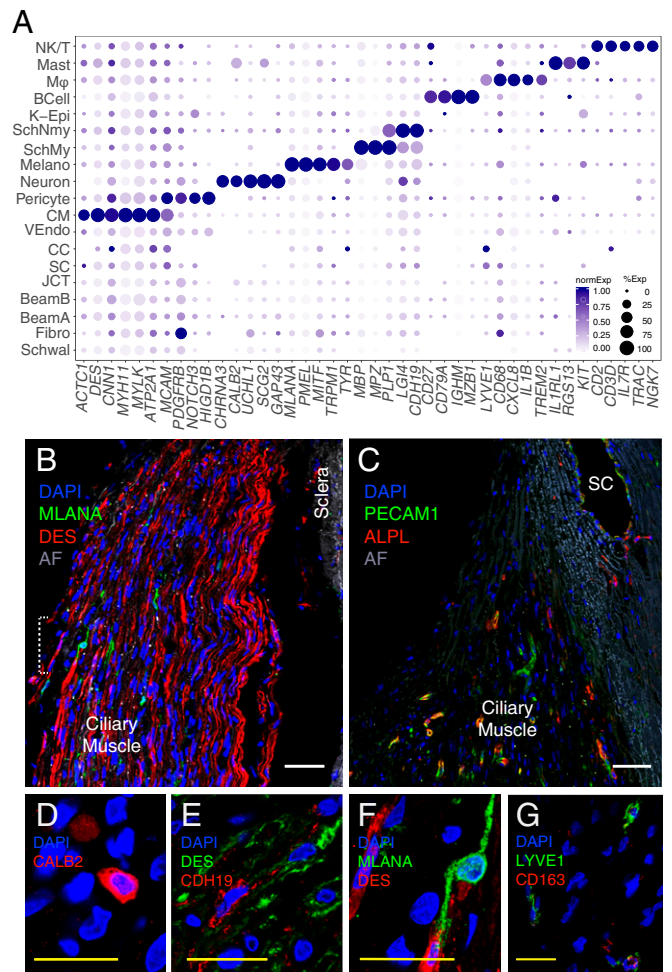


Fig. 4. Cells of the human uveoscleral pathway. (A) Dot plot showing genes selectively expressed in cells of the uveoscleral outflow pathway. Fibro, fibroblast; K-Epi, corneal epithelium; Melano, melanocyte; M ϕ , macrophage; SchMy, myelinating Schwann cell; SchNmy, nonmyelinating Schwann cell; Schwal, Schwalbe line; VEndo, vascular endothelium. (B) Smooth muscle cells immunostained for DES (red) and melanocytes immunostained for MLANA (green) in CM. (C) Capillaries in the CM immunostained for PECAM1 (green) and ALPL (red). Occasional PECAM1+ALPL+ staining was also noted in SC, suggesting that this structure contains more than one cell type. (D) Immunostaining for CALB2 (red) highlights intrinsic neurons of the CM. (E) Immunostaining for LYVE1 (green) and CD27 (red) identifies macrophages in the TM. (F) Schwann cells in the CM immunostained for CDH19 (red) amid CM cells stained for DES (green). (G) Higher magnification of the area bracketed in B demonstrates an MLANA+ melanocyte (green). AF, autofluorescence; DAPI, 4',6-diamidino-2-phenylindole (nuclear stain). (Scale bars: B and C, 50 μ m; D–G, 25 μ m.)

HIGD1B (46). These cells were labeled by immunostaining against *NDUFA4L2* and were wrapped around small vessels (*SI Appendix*, Fig. S2D).

A small neuronal cluster (C19) expressing *CHRNA3*, *UCHL1*, *SCG2*, and *GAP43* was identified and localized to the CM by immunostaining against *CALB2* (aka calretinin) and *ELAVL4* (aka *HuD*) (Fig. 4D and *SI Appendix*, Fig. S2E). These cells presumably correspond to the sparse neurons described in histological and ultrastructural studies (47, 48).

Two clusters, C2 and C17, expressed markers characteristic of Schwann cells, consistent with prior electron microscopic studies showing that Schwann cells ensheath axons within CM (45). Markers common to both clusters included *PLP1* and *LG14*. The clusters were distinguished by differential expression of *CDH19*

in C2 and a set of genes that encode myelin components, such as *MBP*, *MPZ*, and *PMP2*, in C17. Immunostaining for CDH19 marked these cells within the CM (Fig. 4E). We conclude that C2 and C17 represent nonmyelinating and myelinating Schwann cells, respectively.

Finally, C6 comprised uveal melanocytes, demonstrating expression of canonical markers *MLANA*, *PMEL*, *MITF*, *TRPM1*, and *TYR*. Using the marker *MLANA*, we localized these cells to the CM (Fig. 4F).

Immune cells. Our dataset also included four types of immune cells: B cells, Natural Killer (NK)/T cells, mast cells, and macrophages. The macrophages (C4) were *CD163+* and *LYVE1+* (49) and localized predominantly to the TM (Fig. 4G). They also expressed *CD68*, *CD14*, *CCL3*, *CCL4*, *CXCL8*, *IL1B*, *TREM2*, and *MS4A* genes, all of which have been associated with macrophages in other tissues. Mast cells (C13) were localized to the TM using the marker *IL1RL1* and also expressed *CPA3*, *RGS13*, and *KIT* (SI Appendix, Fig. S2F). B cells (C14), characterized by expression of *CD27*, *CD79A*, *IGHM*, *IGKC*, *MZB1*, and *JCHAIN*, were found in only one donor sample but were identified histologically in tissues from other donors using the marker *CD27* (SI Appendix, Fig. S2G). NK/T cells (c10) were identified by differential expression of the genes *CD2*, *CD3D*, *IL7R*, *TRAC*, *GZMA*, *GZMB*, and *NKG7*.

Cell Type-Specific Expression Patterns of Glaucoma-Associated Disease Genes.

To improve understanding of how genes associated with glaucoma contribute to disease pathogenesis, we mapped their expression by the 19 cell types in our atlas. We included both known monogenic causes (Mendelian genes) and genes implicated as risk factors in GWAS (14–22). Mendelian genes assessed were *ANGPT1*, *ANGPT2*, *CPAMD8*, *CYP1B1*, *FOXC1*, *LOXL1*, *LTBP2*, *MYOC*, *OPTN*, *PITX2*, *TEK* (*TIE2*), and *TBK1*. Well-established GWAS loci genes included *TMCO1*, *TXNRD2*, *CAV1*, *CAV2*, *ARHGEF12*, and *ATXN2*. We also screened an additional 462 genes listed in the National Human Genome Research Institute–European Bioinformatics Institute (NHGRI-EBI) GWAS Catalog for notable expression patterns (50). Overall, 189 genes were expressed in a minimum of 10% of cells in any given class at a minimum expression level of 0.5. Examples are shown in Fig. 5, with a full list in SI Appendix, Fig. S3.

Most genes implicated in juvenile glaucoma and high IOP (e.g., *MYOC*, *FOXC1*, *PITX2*, *CYP1B1*) were expressed strongly by all three TM cell types (Beam A, Beam B, and JCT), with no evidence for selective expression by any one of the three (Fig. 5 and SI Appendix, Fig. S2 K and L). Notably, they were all also expressed at high levels in CM. *EFEMP1* exhibited robust cell type-specific expression in beam cells and JCT; in contrast to the first group, it was not present in CM and was expressed differentially among the TM types: JCT > Beam B > Beam A. In contrast, some genes demonstrated stronger expression in non-TM cell types than in TM, including *GAS7* (CM, Schwann cells), *KALRN* (CM), *PRSS23* (SC, CC, vessel endothelium, and Schwann cells), and *CAV1/CAV2* (all three endothelial types, CM, pericytes, and melanocytes). Thus, although defects in the TM are clear contributors to elevated IOP, genes that regulate IOP may not act exclusively within the TM.

Because glaucoma risk involves susceptibility of RGCs to degeneration as well as increased IOP, we also interrogated expression in four relevant retinal cell types: RGCs and three glial types with which they interact (astrocytes, Müller glia, and microglia). Data for retinal cells are taken from a human retinal cell atlas (*Materials and Methods*) (23). We found a distinction between genes associated with primary open-angle glaucoma (POAG) involving elevated IOP and those for which an IOP-independent component may exist. Whereas most genes associated with POAG and/or elevated IOP were expressed in cells of

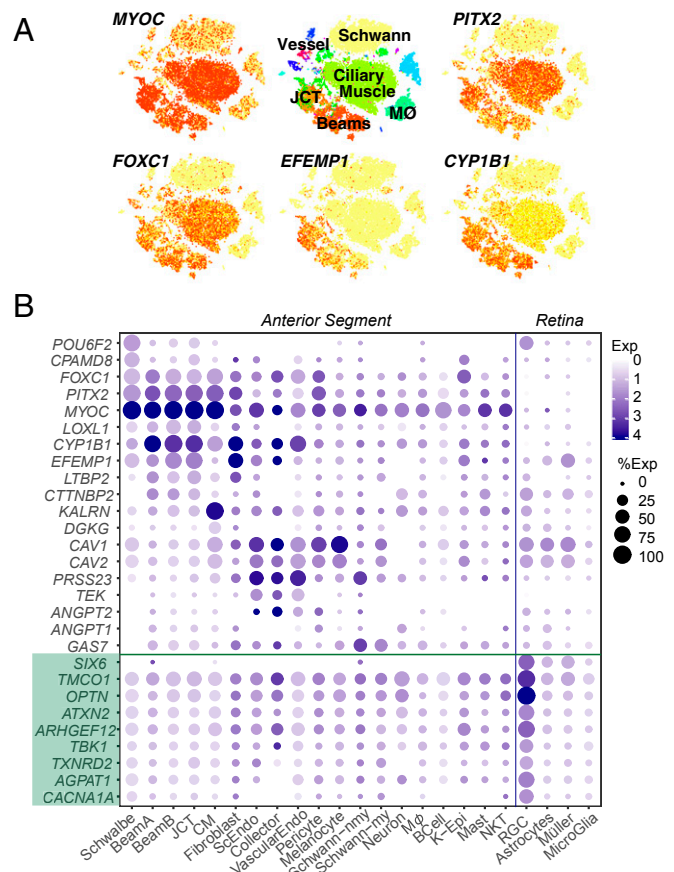


Fig. 5. Human disease genes. Cell type-specific expression of several genes implicated in glaucoma as illustrated by t-distributed stochastic neighbor embedding (A; arranged as in Fig. 2A) and a dot plot (B). B also includes data from a retinal cell atlas, showing expression in RGCs and three types of retinal glia. Green shading highlights genes that genetics studies suggest may confer additional IOP-independent glaucoma risk; many of these genes are also associated with high IOP. K-Epi, corneal epithelium; MØ, macrophage; SEndo, Schlemm canal endothelium; Schwann-my, myelinating Schwann cell; Schwann-nmy, nonmyelinating Schwann cell; VascularEndo, vascular endothelium.

the outflow pathways as discussed above, a subset, including *OPTN*, *ATXN2*, *TMCO1*, and *SIX6*, were expressed at the highest levels in RGCs (Fig. 5B).

Several susceptibility genes, including *CAV1*, *CAV2*, and *POU6F2*, were expressed at high levels in both the anterior segment and retina. *POU6F2* is a transcription factor that has been linked to thinner than average central corneal thickness, a highly heritable trait and also a strong risk factor for the development of POAG (51). The expression of *POU6F2* has been detected in human RGCs (52), mouse corneal limbal stem cells, and a subset of mouse RGCs (53). We found preferential expression of *POU6F2* in both corneal endothelium (specifically, Schwalbe line cells) and RGCs (Fig. 5B).

Two genes associated with exfoliation glaucoma (XFG) and the related exfoliation syndrome (XFS), *LOXL1* and *CACNA1A*, demonstrated divergent regional expression patterns: the former localized predominantly to TM (beam cells and JCT), whereas the latter localized preferentially to RGCs. This result, consistent with prior investigations (22), offers further insight into XFG as both a primary and a secondary open-angle glaucoma characterized not only by IOP elevation due to TM obstruction but also by an inherent susceptibility to glaucomatous optic neuropathy.

Finally, motivated by literature implicating the complement system, aquaporins, matrix metalloproteinases (MMPs), cytochrome P450 enzymes, and prostaglandins in ocular disease pathogenesis, progression, and treatment (54, 55), we investigated their expression in the anterior segment. We found that many genes encoding complement factors were selectively expressed in the conventional outflow pathway. For example, expression of *CIQA*, *CIQB*, and *CIQC* was observed in resident macrophages, whereas the serine proteases *CIR* and *CIS*; the C1 inhibitor *SERPING1*; and other complement genes *CFD*, *CFH*, *C3*, and *C7* were expressed preferentially in both TM cells and scleral fibroblasts (*SI Appendix*, Fig. S4A). Among MMPs, *MMP2* and *MMP14* were most predominant (*SI Appendix*, Fig. S4C); the former was selectively expressed by beam cells and SC endothelium as well as scleral fibroblasts, whereas the latter was expressed mostly in scleral fibroblasts. *AQPI* and *CYP1B1* were uniquely expressed in outflow cell types among their respective family members (*SI Appendix*, Fig. S4B and E). Within the Rho-protein family, *RHOB* demonstrated the strongest expression among outflow cell types followed by *RHOA* and *RHOC* (*SI Appendix*, Fig. S4D). The prostaglandin genes *PTGDS* and *PTGES3* were robustly expressed in all outflow cell types, whereas *PTGS2*, *PTGRI*, and *PTGER2* were more selectively expressed in the conventional outflow pathway cells (*SI Appendix*, Fig. S4F).

Model Species. Analysis of mechanisms underlying IOP regulation and preclinical studies of therapeutic interventions rely almost entirely on model species. Yet, limited information is available on how closely cells and molecules of the anterior segment in commonly used models resemble those of humans. To address this issue, we profiled cells dissociated from anterior segment tissues of four model species: the rhesus macaque (*M. mulatta*; 5,158 cells), the cynomolgus macaque (*M. fascicularis*; 9,155 cells), the common swine (*S. scrofa*; 6,709 cells), and mouse (*M. musculus*; 5,067 cells). These species are among the most commonly used for studies on glaucoma: rhesus and cynomolgus monkeys are frequently used for basic studies of primate visual physiology and preclinical tests, respectively (26); porcine anterior segments are used in aqueous outflow studies (24); and the broadest range of genetically modified lines is available in mice (25). We used a machine learning algorithm [XGBoost (56)] (*Materials and Methods*) to find correspondences of cell types in the model species with those in humans. While significant conservation was noted among certain cell types across species, there were also notable differences.

Macaca mulatta. Cells collected from anterior chamber angle structures of *M. mulatta* clustered into 15 types (Fig. 6A and E). We identified three TM-like clusters in this species, similar to humans (Fig. 6B). One cluster corresponded to human Beam Cell A (mmC4), sharing markers including *BMP5* and *EDN3*; another corresponded to human JCT (mmC10), sharing preferential expression of markers including *ANGPTL7* and *CHI3L1*. The third cluster (mmC1), which we call Beam Cell X, could not be mapped to a single human cluster but instead, shared multiple markers of all human TM types; it was distinguished from mmC4 and mmC10 by its preferential expression of *CYP1B1* and *MGARP*. We identified types mapping 1:1 to human SC (mmC7), vascular endothelium (mmC8), and the uveoscleral outflow pathway, including CM (mmC2), melanocytes (mmC9), nonmyelinating Schwann cells (mmC5), and myelinating Schwann cells (mmC14). Two types of pericytes were present (mmC11, mmC12) compared with one in humans. Among immune cells, macrophage (mmC3, mmC15) and NK/T (mmC6, mmC13) clusters were present. B cells, mast cells, and neurons were not detected, possibly due to their sparsity. Finally, due to a conservative dissection of the TM strip in this species, CC endothelium, corneal epithelium, and Schwalbe line cells were not recovered.

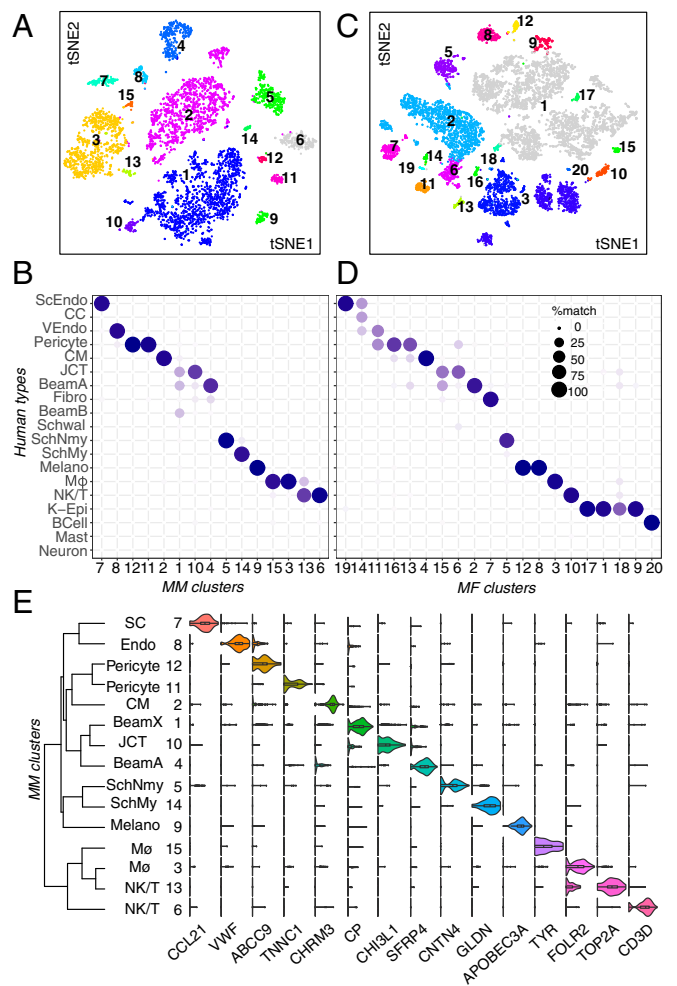


Fig. 6. Cell types and gene expression in the outflow pathways of two macaque species (*M. mulatta*, MM; and *M. fascicularis*, MF). (A) A t-distributed stochastic neighbor embedding (tSNE) plot showing 15 cell types derived from TM and associated structures of *M. mulatta*. (B) Transcriptional correspondence between human and *M. mulatta* cell types summarized as a “confusion matrix.” In this and subsequent figures, the size of the circle and its intensity indicate the percentage of cells of a given cluster from the model species (column) assigned to a corresponding human cluster (row) by a classification algorithm trained on the human cells. Fibro, fibroblast; K-Epi, corneal epithelium; Melano, melanocyte; MØ, macrophage; VEndo, vascular endothelium; SchMy, myelinating Schwann cell; SchNmy, nonmyelinating Schwann cell; Schwal, Schwalbe line; VEndo, vascular endothelium. (C) A tSNE plot showing 20 cell types derived from TM and associated structures of *M. fascicularis*. (D) Transcriptional correspondence between human and *M. fascicularis* shown as in B. (E) Violin plot showing examples of genes selectively expressed by each cell type in *M. mulatta*.

Macaca fascicularis. Cells collected from anterior chamber angle structures of *M. fascicularis* clustered into 19 types (Fig. 6C and *SI Appendix*, Fig. S5A). We identified three TM-like clusters (mfC2, mfC6, mfC15). Of these, one cluster, mfC2, preferentially expressed *BMP5* and *EDN3* and could be confidently mapped to human Beam Cell A (Fig. 6D). The other two clusters (mfC6 and mfC15) expressed genes observed in multiple human TM clusters, including *MYOC*, *MGP*, and *ANGPTL7*; overall, mfC6 shared more genes with the human JCT cluster (e.g., *FMOD*, *CEMIP*) whereas mfC15 expressed a combination of genes found within the the human beam cell, fibroblast, and Schwalbe line clusters (e.g., *ANGPTL5*, *COL6A3*, *AQPI*, and *POU3F3*) (*SI Appendix*, Fig. S5A). Three vascular endothelial clusters were identified (mfC11, mfC14, mf19); these corresponded

to the human capillary endothelium, CCs, and SC endothelium, respectively. Other clusters mapped closely to cell types in the human dataset, including CM (mfC4), Schwann cells (mfC5), macrophages (mfC3), B cells (mfC20) and NK/T cells (mfC10), melanocytes (mfC8, mfC12), and pericytes (mfC13, mfC16). No neurons were identified.

Based on results from human, we profiled cells from one sample that included the residual corneal scleral rim in addition to TM strips in hopes of identifying CC endothelium and matrix fibroblasts. Indeed, both additional cell types were recovered in addition to multiple epithelial cell types (not described here). Matrix fibroblasts appeared during the initial unsupervised clustering as mfC7. CC endothelium (mf14) could be identified within the original SC cluster using a supervised approach based on differential expression patterns identified in humans (*SI Appendix, Fig. S5B*).

Sus scrofa. Molecular profiling of porcine anterior segment tissue yielded 18 clusters (Fig. 7A). Two clusters comprised TM cells. One mapped to human Beam Cell A (ssC1), with differential expression of *SFRP4* and *TMEFF2* in addition to *PON1*, *FMO1*, and *RARRES1*. The other mapped to human JCT (ssC3), selectively expressing *CCN3*, *FMO3*, *CEMIP*, and *BMP3* (Fig. 7B and C). While expression of *MYOC* was low across all clusters, it was preferentially expressed in ssC1. Both TM clusters expressed *CYP11B1*, *MGP*, *CFH*, and *NR2F1*.

Of two endothelial clusters identified, one mapped to human SC (ssC14), and the other mapped to capillary endothelium (ssC12). While the pig does not possess a continuous SC and instead, has more discontinuous vessels referred to as an aqueous plexus, cells lining this structure resemble those of human SC. The SC was distinguished from capillary endothelium through differential expression of lymphatic markers *CCL21*, *PROX1*, and *LYVE1* and was histologically validated with immunostaining against *LYVE1* (*SI Appendix, Fig. S6A*).

A CM cluster (ssC16) was marked by preferential expression of *DES* and *ACTA2*, both of which we confirmed with immunostaining; the latter also stained pericytes (*SI Appendix, Fig. S6B*). *NDUFA4L2* was expressed predominantly by pericytes (ssC11) and CM but also, corneal endothelium and to a lesser degree, TM cells (*SI Appendix, Fig. S6C*). The low yield of CM in our dissection can be attributed to the porcine angle anatomy, which differs from that of primates in having no clear anatomical connection between the CM and conventional drainage structures (57). It is, therefore, easily excluded during dissection.

Other clusters corresponding to human cell types included myelinating (ssC18) and nonmyelinating (ssC13) Schwann cells, melanocytes (ssC15), fibroblasts (ssC6), corneal endothelium (ssC17), macrophages (ssC4), and NK/T cells (ssC10). Five clusters (ssC2, -5, -7 to -9) were derived from ocular surface epithelium due to a permissive dissection technique in which the TM/corneal scleral rim was dissociated. Cell types absent in this collection included mast cells, B cells, and neuronal types.

Mus musculus. The mouse eye is small compared with those of other species profiled, and therefore, we dissociated the entire anterior segment, including cornea, iris, and ciliary body. Presumably for this reason, initial clustering by cell class signatures revealed that 8,766 of 13,833 single-cell profiles corresponded cells of the ocular surface epithelium. We eliminated these cells and reanalyzed the remaining 5,067 cells, yielding 20 clusters (Fig. 8A and C).

Although the mouse iridocorneal angle is more compact than that of humans, anatomical studies suggest that conventional outflow structures in mice are similar in many respects, with trabecular lamellae, distinctive JCT, and a continuous SC (58). We identified three candidate TM clusters (mC6, mC9, and mC14) through their high levels of *Myoc* expression. Two of these clusters, mC14 and mC6, mapped preferentially to human Beam Cell A and JCT, respectively, while also demonstrating some ECM expression patterns similar to the human corneal

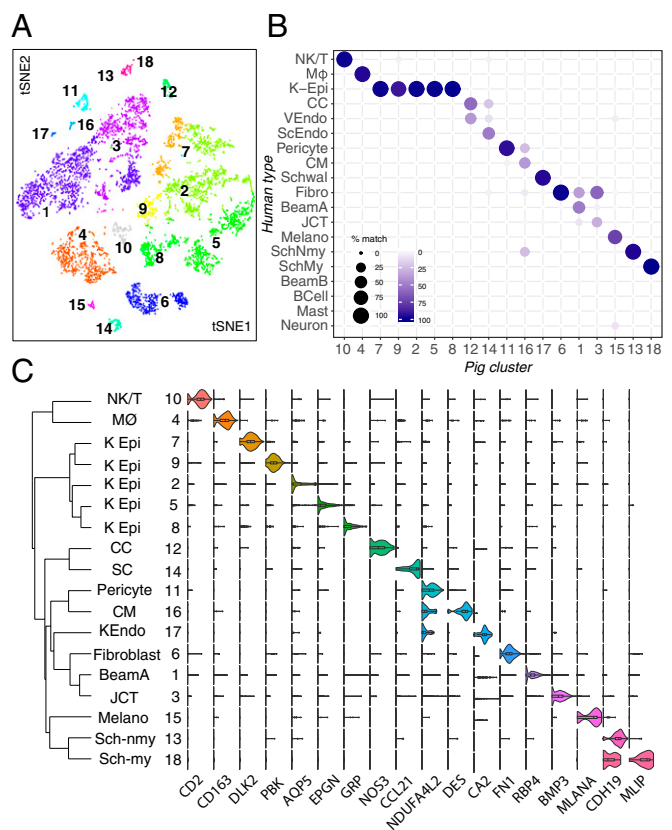


Fig. 7. Cell types and gene expression in the outflow pathways of the pig. (A) A t-distributed stochastic neighbor embedding (tSNE) plot showing 18 cell types derived from TM and associated structures of pig. (B) Transcriptional correspondence between human and pig cell types shown as in Fig. 6B. Fibro, fibroblast; K-Epi, corneal epithelium; Melano, melanocyte; MΦ, macrophage; SchMy, myelinating Schwann cell; SchNmy, nonmyelinating Schwann cell; Schwal, Schwalbe line; VEndo, vascular endothelium. (C) Violin plot showing examples of genes selectively expressed by each cell type in pig. KEndo, corneal endothelium; K Epi, corneal epithelium; ScEndo, Schlemm canal endothelium; Sch-my, myelinating Schwann cell; Sch-nmy, nonmyelinating Schwann cell.

fibroblast cluster (Fig. 8B). The third candidate TM cluster, mC9, despite sharing many markers with fibroblasts (e.g., *Pi16*, *Fbn1*, *Mfap5*, *Tnxb*, *Clec3b*), was closely related to mC14 and thus, tentatively named Beam Cell Y. All three TM clusters also expressed *Mgp* and *Pdpr* similar to human TM cells; the latter was confirmed histologically with immunostaining as was expression of *Chil1* (*Chi3l1*) in mC6 and mC14 (Fig. 8D and E and *SI Appendix, Fig. S7*). Other differentially expressed genes within the mouse JCT cluster included *Nell2*, *Chad*, and *Tnmd*, whereas differentially expressed genes within the mouse beam clusters included *Sfp4* (both mC9 and mC14), *Tmeff2* (mC9), and *Fmo2* (mC14).

Among other outflow cell types, we identified two separate vessel clusters (mC10, mC19) corresponding to vascular and SC endothelium, respectively. *Postn*, which was also differentially expressed within the human SC cluster, was also expressed in mouse SC as well as JCT (Fig. 8F and G).

Other clusters mapping to human cell types included corneal endothelium/Schwalbe line (mC12), smooth muscle (mC17), pericytes (mC8), Schwann cells (mC18), uveal melanocytes (mC2), macrophages (mC4), B cells (mC20), and NK/T cells (mC16) (Fig. 8B). In contrast to our human data, the smooth muscle and Schwann cell clusters were small, which can be attributed in large part to the diminutive murine accommodative apparatus (58).

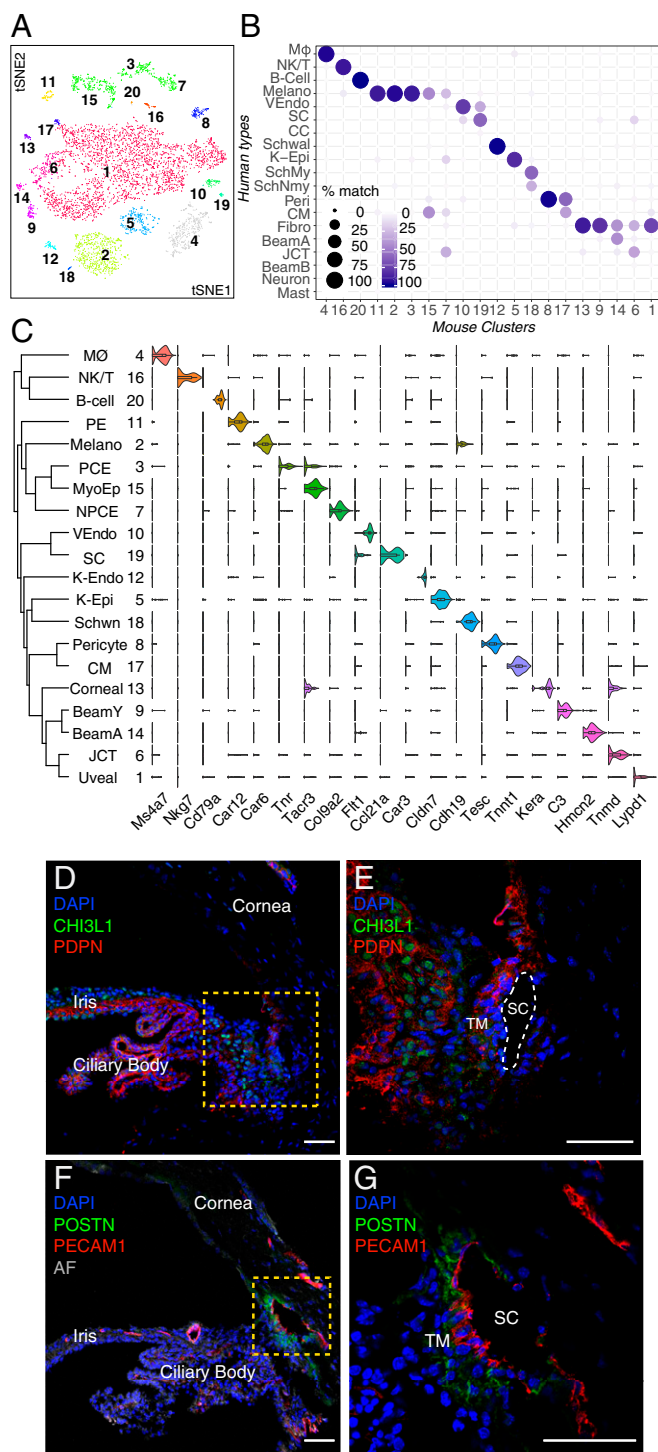


Fig. 8. Cell types and gene expression in the outflow pathways of the mouse. (A) A t-distributed stochastic neighbor embedding (tSNE) plot showing 20 cell types derived from TM and associated structure of mouse. (B) Transcriptional correspondence between human and mouse cell types shown as in Fig. 6B. Fibro, fibroblast; K-Epi, corneal epithelium; Melano, melanocyte; Mφ, macrophage; SchMy, myelinating Schwann cell; SchNmy, nonmyelinating Schwann cell; Schwab, Schwalbe line; VEndo, vascular endothelium. (C) Violin plot showing examples of genes selectively expressed by each cell type in mouse. K-Endo, corneal endothelium; NPCE, nonpigmented ciliary epithelium; PE, pigmented epithelium; SchwN, Schwann cell. (D and E) Pdpn (red) is present in multiple cell types, including pigmented and nonpigmented epithelium of the iris and ciliary body (CB) as well as a subset of TM cells, K-Endo, and K-Epi, whereas Chil1 (green; ortholog to CHI3L1 in humans) stains a different subset of TM cells and to a

Owing to the dissection method, the mouse dataset also included a large number of cells derived from iris and ciliary body. We tentatively identified these clusters based on markers identified in previous studies (43, 59); they comprised melanocytes, stromal fibroblasts, pigmented epithelium, and nonpigmented epithelium of the iris and ciliary body.

Conservation of Expression Patterns among Species. Next, we assessed expression patterns in model species of key genes selectively expressed in cell types of the human AH outflow pathways. In general, conservation was striking.

Many markers expressed across all human TM clusters were conserved in other species (Fig. 9A). They included matrix-related genes, such as *DCN* and *MGP*, and the retinoic acid-related genes *RARRES1* and *RBP*. However, some differences among species were evident. *PDPN*, a selective marker for TM cells in humans, was less selective for these cells in other species. For example, it was also found at similar levels in uveal melanocytes in all four nonhuman species. *BMP5*, expressed in Beam Cell A in human and macaque, was present but less specific in pig and absent in mouse. The human JCT marker, *CHI3L1*, was preferentially expressed in JCT clusters in monkey and mouse but was notably absent in pig.

There was also excellent correspondence between markers of other cell types that compose the conventional pathway. Markers of CC, SC, and vascular endothelium markers were generally well conserved among human and macaque (SI Appendix, Fig. S8A). Pig and mouse also shared many markers; however, SC in these species tended to demonstrate more prominent expression of lymphatic markers than primates. Among species in which Schwalbe line/corneal endothelial cells were obtained, there was also good correspondence (SI Appendix, Fig. S8B). Of note, whereas the Schwalbe line cluster in humans includes cells at the junction between cornea and TM, the corresponding clusters in pig and mouse more likely represent a larger proportion of corneal endothelia. Markers of cell types comprising the uveoscleral pathway and immune cells were also very well conserved among species (Fig. 9B and SI Appendix, Fig. S8C).

Finally, we analyzed patterns of disease gene expression in these model species (Fig. 9C). To facilitate comparison, individual TM cell clusters from each species were merged during this analysis, and a threshold was set such that only genes with expression in >10% of cells in at least one cluster were plotted. While some genes were consistently expressed in corresponding cell types across species, others exhibited significant differences. For example, we noted that the classic anterior segment dysgenesis genes *FOXC1* and *PITX2* were reliably expressed in TM cell types across all species. Similarly, *MYOC* was also expressed in TM cells across species; however, in this case, subtle differences in other cell types were observed. For example, in human and macaque, *MYOC* was strongly expressed in both TM and CM, whereas in pig and mouse, it was expressed only in TM. Similarly, *MYOC* was highly expressed in Schwalbe line cells of human but not in the corresponding corneal endothelial clusters of pig (ssC17) and mouse (ssC12). As mentioned above, this is likely because ssC17 and mC12 represent a larger set of corneal endothelia, not just the peripheral subset. *LOXLI*, while expressed in TM clusters of human, pig, and mouse, was absent in the TM of both macaque species. *KALRN*, expressed in human, macaque, and mouse CM, was absent in pig. Finally, *CYP11B1*, expressed in TM cells across

lesser extent, cells within the CB. (E) Higher magnification of the boxed area in D. (F and G) Immunostaining against Postn (green), a secreted protein, highlights SC and JCT cells. Pecam1 (red) highlights SC as well as vascular endothelial clusters. AF, autofluorescence. DAPI, 4',6'-diamidino-2-phenylindole (nuclear stain). (Scale bars: 50 μm.)

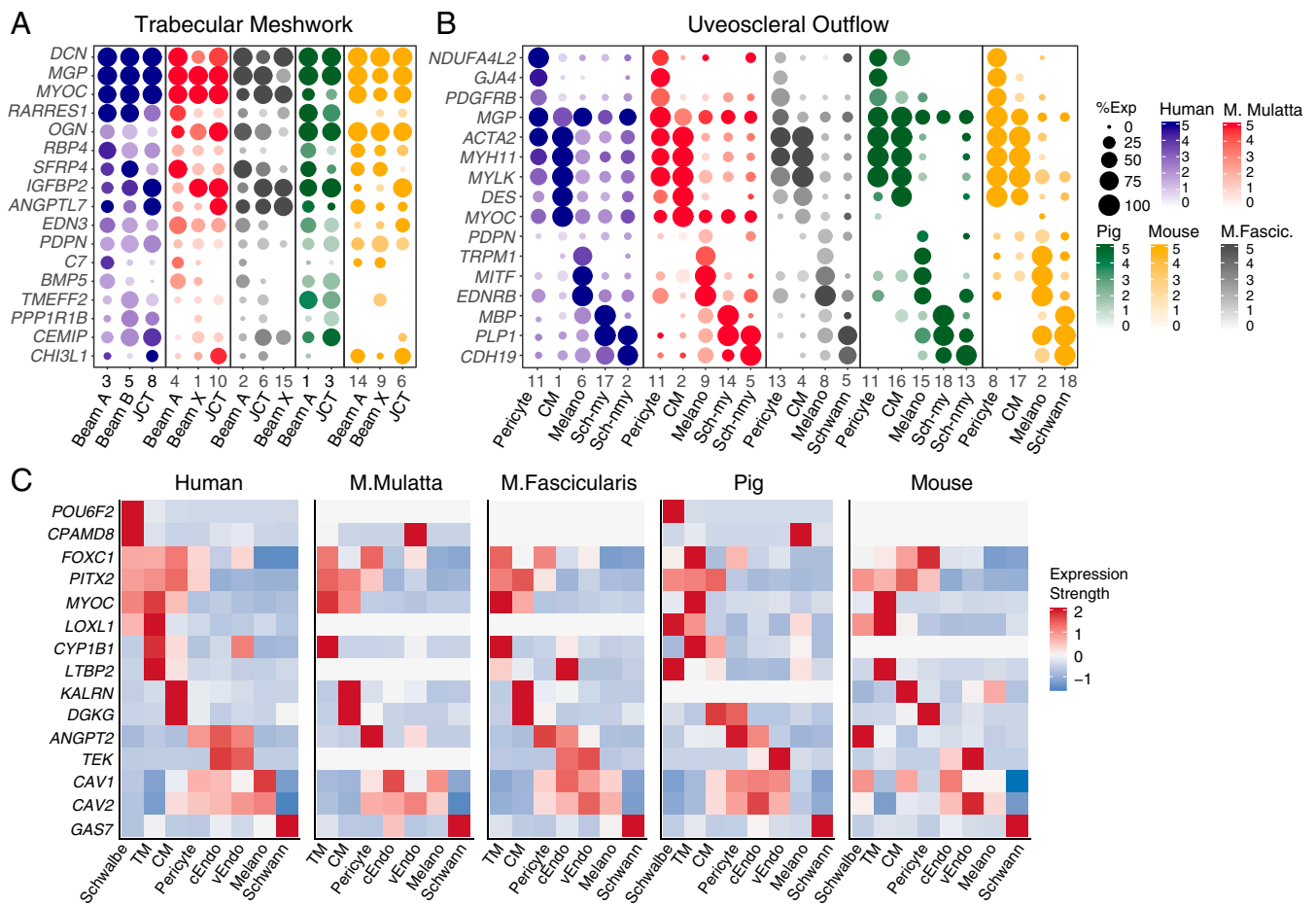


Fig. 9. Comparison of gene expression across species. (A and B) Key genes are shown in dot plots for cell types comprising the TM (A) and uveoscleral outflow pathway (B). (C) Heat map showing expression of genes implicated in human POAG in aqueous outflow cells of humans (replotted from *SI Appendix, Fig. S3*) and model species. In some cases (i.e. TM, cEndo), multiple cell types from each species have been merged into their broader class to facilitate comparison. cEndo, canal endothelium (includes both Schlemm canal and collector channel clusters where available); Melano, melanocyte; Sch-my, myelinating Schwann cell; Sch-nmy, nonmyelinating Schwann cell; TM, trabecular meshwork (includes beam cells and JCT where available), vEndo, vascular endothelium.

human, macaque, and pig, did not meet threshold expression levels in mouse TM. This is consistent with prior studies, which have reported *CYP1B1* complementary DNA but not protein expression in human adult TM samples and negative immunohistochemistry (IHC) staining in mouse TM (60).

Discussion

We used scRNAseq to profile cells composing the AH outflow pathways in humans, generating a cell atlas for these tissues and identifying markers for each cell type. We then used the atlas to localize the expression of genes implicated in glaucoma. These findings offer insights into the molecular architecture of TM cells and highlight potential roles in IOP homeostasis for non-TM cell types in the anterior chamber angle. Finally, we profiled cells in the outflow pathways of four model species—*M. mulatta*, *M. fascicularis*, *S. scrofa*, and *M. musculus*—providing a foundation for using these models in studies on regulation and dysregulation of IOP.

Technical Issues. A major challenge in human vision research is that essentially all ocular tissues must be obtained either post-mortem or postnucleation. Furthermore, postmortem ocular tissues of the anterior segment suitable for transplantation are understandably prioritized for this purpose over research. In this study, all but one of the postmortem tissues sequenced were obtained within ~6 h of death from a rapid autopsy program

enrolling predominantly oncology patients. While donors had no documented pathological ocular history or clinical evidence of eye disease on examination by an ophthalmologist (T.v.Z), they did have varying degrees of chronic systemic disease and different levels of antemortem exposure to chemotherapeutic and steroid medications. This represents a limitation of our study. Furthermore, lack of documented or physically evident ocular pathology cannot be taken as evidence that none existed. Since we focused primarily on cell type classification in this study rather than quantitative determination of gene expression levels, we believe that the donors' systemic diseases did not influence the ultimate cell atlas. This was further corroborated in two ways: 1) demonstrating that the same cell types were obtained from all individuals and 2) performing histological validation on tissues obtained from a wider variety of donors through an eye bank as well as on separate rapid autopsy patients.

Human Cell Atlas. In the human AH outflow pathways, we identified 19 major cell types: 8 belonging to the conventional outflow pathway (including 3 distinct populations within the filtering TM), 7 belonging to the uveoscleral pathway, and 4 representing immune cell populations.

By histological criteria, the filtering TM has been divided into three layers: a uveal layer adjacent to the anterior chamber, a juxtacanalicular layer adjacent to SC, and a corneoscleral layer in

between (7, 9). The JCT cells in our dataset clearly localized closest to SC, with the Beam A and B cells localizing to the other two layers. It is tempting to assign Beam A and B to uveal and corneoscleral layers, respectively, but our histological analysis suggests that they are in fact intermingled.

The lymphatic marker podoplanin (*PDPN/D2-40*) emerged as a robust marker for all human TM cell types (C3, C5, and C8), consistent with previous studies (31, 32). Along with *CCL2* (a chemoattractant for monocytes) and *VCAM1* (a mediator of immune cell migration), there was a marked specificity of *PDPN* expression among cells of the conventional pathway compared with the uveoscleral pathway, suggesting that the former acts as an immunological “sink” guiding antigen presenting cells and other immune cells toward SC, the venous system, and ultimately, the spleen (36). Consistent with this idea, SC expresses markers of lymphatic vessels (e.g., *CCL21* and *FLT4/VEGFR3*), whereas CCs express venular markers (e.g., *ACKR1*).

Model Species. In comparing the cellular composition of human AH outflow pathways with those of commonly used models, we documented excellent correspondence for many cell types across species. CM cells, pericytes, melanocytes, and Schwann cells were present in all species and demonstrated shared expression of canonical markers. Corneal endothelium was identified in pig and mouse but in neither macaque species, likely due to our targeted dissection technique. Two or more types of vascular endothelium, defined as *PECAM1 + TIE1+* clusters, were also present in each species and could be assigned either to SC or to other vasculature. Among immune cells, *LYVE1 + CD163 + CD68+* macrophages were also present in all species; other immune cells were identified in each species with less correlation, perhaps due to their relatively sparse numbers. Similarly, neurons were limited to the human dataset, most likely due to their low numbers.

On the other hand, while TM cells could be identified in each species, they demonstrated substantial variability across species. Three TM types—two beam cell types and one JCT cell type—were present in all three primates as well as the mouse, but they failed to map 1:1 across species. Beam A and JCT cells were present in these four species, but the second beam type, which we call Beam X in both macaque species and Beam Y in the mouse, demonstrated more unique expression patterns. Moreover, we identified only one beam cell type in pig, although it is possible that further analysis with increased sample size would allow a subdivision of this cluster. Some differentially expressed markers, such as *PDPN*, *RARRES1*, *CH3L1*, and *ANGPTL7*, which were selectively expressed by beam and/or JCT cells in human tissue, were either absent in the model species or nonspecific. Others were conserved across species, including *MYOC*, *EDN3*, and *RBP4*. Interestingly, *PDPN* did not demonstrate the same specificity in any of our model species, suggesting that this may be a human-specific feature.

Glaucoma. Glaucoma is a phenotypically heterogeneous disease with traits dictated by complex interactions among age, environment, and genes. Examination of cell type-specific expression patterns of both Mendelian genes and GWAS susceptibility loci revealed multiple expression patterns, of which we note four groups. First, several IOP-associated disease genes were selectively expressed by TM cell types (e.g., *CYP11B1* and *EFEMP1*), supporting their potential contribution to function and or/dysfunction of this tissue. Second, others were expressed not only by TM but also by cells of the uveoscleral pathway (e.g., *MYOC*, *PITX2*, and *FOXCI*). Third, a few genes implicated in high IOP mapped selectively to cells of the uveoscleral pathway (e.g., *KALRN*). Together, these results may indicate an integral contribution of the uveoscleral pathway to disorders of IOP. Fourth, some genes were expressed by RGCs, often in addition to TM cells. For example, *TMCO1*, while clearly expressed in the

anterior segment outflow pathways, also demonstrated robust expression in RGCs, supporting the notion that it may also confer risk for glaucoma development through an IOP-independent mechanism (61).

Although TM cells have been shown to exhibit phagocytic abilities, it is possible that resident macrophages in the conventional outflow pathway also contribute to the phagocytic workload. Their stimulation or recruitment to the TM may serve to lower IOP as has been suggested to occur after selective laser trabeculoplasty (62). Conversely, dysfunctional macrophages may contribute to elevated IOP (63).

Conclusion

In conclusion, we have generated a cell atlas of the trabecular meshwork and surrounding structures constituting the aqueous humor outflow pathways in humans and four model species. We identified markers for each cell type, enabling in-depth study of individual components of this complex tissue and shedding light on features such as the immunological milieu and the lymphatic characteristics of the conventional outflow pathway. Our atlas also provides biological context for susceptibility loci identified in genome-wide association studies. Together, these results may guide further investigation in both humans and model species.

Materials and Methods

Tissue Acquisition and Processing. Human ocular tissues used for sequencing, immunohistochemistry, and in situ hybridization were collected a median of 6 h postmortem (range 3 to 14 h) (*SI Appendix, Table S1*). Human tissue used for IHC and in situ hybridization was provided by the Lions Vision Gift, Portland, OR. No ocular disease was reported in any of the human donors, and no abnormalities were noted during microdissection. Nonhuman primate eyes were obtained from macaques 4 to 10 y of age that had reached the end of unrelated studies at supplying institutions. Porcine (*S. scrofa*) eyes were obtained from a local abattoir. Murine eyes were collected from male and female 12-wk-old CD1 mice obtained from Charles River Laboratories.

Procedures were conducted in compliance with the Association for Research in Vision and Ophthalmology’s Statement for the Use of Animals in Ophthalmic and Vision Research and the guidelines for the care and use of animals and human subjects at Harvard University and Partners Healthcare. Acquisition and use of human tissue were approved by the Human Study Subject Committees (Dana Farber/Harvard Cancer Center Protocol No. 13–416 and Massachusetts Eye and Ear - Non-Human Subjects Research Protocol No. 18–034H). Acquisition and use of nonhuman tissue were approved by the Institutional Animal Care and Use Committee at Harvard University.

RNA Sequencing. Tissues were digested enzymatically, and single cells were loaded into 10× Chromium Single Cell Chips (27). Single-cell libraries were generated following the manufacturer’s protocol. Libraries were sequenced on Illumina HiSeq 2500. Sequencing data were demultiplexed and aligned using Cellranger software (10× Genomics). Reads were aligned to the following reference genome: human samples, GRCh38; *M. mulatta*, Mmul8; *M. fascicularis*, MacFas5 with our augmented transcriptome file (64); pig, Sscrofa11; and mouse, mm10. Subsequent clustering and identification of differentially expressed genes followed methods described by Peng et al. (64) with modifications detailed in *SI Appendix, Supplementary Materials and Methods*. Transcriptomic similarity between humans and other model species was assessed with a machine learning algorithm XGBoost (56).

Histology. Corneoscleral wedges were fixed, embedded in tissue freezing medium, and sectioned in a cryostat. *SI Appendix, Table S2* shows donor information. Sections were immunostained or processed for in situ hybridization using probes listed in *SI Appendix, Table S3*. Images were acquired on Zeiss LSM 710 confocal microscopes.

Data Availability. The accession number for the raw and unprocessed data files reported in this paper is Gene Expression Omnibus accession number GSE146188. Data can be visualized in the Broad Institute’s Single Cell Portal at https://singlecell.broadinstitute.org/single_cell/study/SCP780.

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