Quantitative single-cell transcriptome-based ranking of engineered AAVs in human retinal explants

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Gene therapy is a rapidly developing field, and adeno-associated viruses (AAVs) are a leading viral-vector candidate for therapeutic gene delivery. Newly engineered AAVs with improved abilities are now entering the clinic. It has proven challenging, however, to predict the translational potential of gene therapies developed in animal models due to cross-species differences. Human retinal explants are the only available model of fully developed human retinal tissue and are thus important for the validation of candidate AAV vectors. In this study, we evaluated 18 wild-type and engineered AAV capsids in human retinal explants using a recently developed single-cell RNA sequencing (RNA-seq) AAV engineering pipeline (scAAVengr). Human retinal explants retained the same major cell types as fresh retina, with similar expression of cell-specific markers except for a photoreceptor population with altered expression of photoreceptor-specific genes. The efficiency and tropism of AAVs in human explants were quantified with single-cell resolution. The top-performing serotypes, K91, K912, and 7m8, were further validated in non-human primate and human retinal explants. Together, this study provides detailed information about the transcriptome profiles of retinal explants and quantifies the infectivity of leading AAV serotypes in human retina, accelerating the translation of retinal gene therapies to the clinic.

INTRODUCTION

The FDA approval of adeno-associated viruses (AAVs) for treatment of Leber congenital amaurosis is a milestone in the field of gene therapy,1 and a variety of AAV-mediated gene therapies for a wide range of diseases such as inherited retinal dystrophies, neuromuscular disorders, hemophilia, and inherited metabolic disorders are at an advanced stage of clinical development.2,3 Efficient gene delivery is key to the success of each of these gene therapy approaches. Levels of transgene expression are determined by the properties of the viral capsid, the promoter driving transgene expression, the injection route, and the time point of delivery.4 A variety of studies have shown that changes can be made to the structure of the protein capsid shell of the virus and that reengineering of the viral capsid can result in AAVs with improved transduction abilities, enabling the application of lower doses of AAV, and diminished risks of adverse effects.5 Mutation of surface-exposed tyrosine, serine, and threonine residues resulted in viruses with increased infectivity and protection of the AAV capsid from degradation. And, directed evolution (DE) approaches have resulted in the development of newly engineered AAVs such as 7m8, which has an improved ability to bypass structural barriers in the retina and infect photoreceptors.6 A number of clinical trials using these next-generation engineered viral vectors are ongoing (ClinicalTrials.gov: NCT03316560, NCT02416622, and NCT03748784).

It is essential to accurately quantify transgene expression levels and cell-type tropism of vectors to accurately predict the success of gene therapies in patients. Mice have, to date, been the mostly widely used preclinical animal model for such studies. However, due to anatomical and structural differences, AAV expression patterns differ dramatically in the retina of mouse and non-human primates (NHPs), the animal model with the highest similarity to humans.6 The quantification and validation of AAVs in NHPs has, in the past, been difficult due to variability between animals, as well as the cost and ethical burdens associated with such work. Therefore, we recently established a single-cell RNA sequencing (RNA-seq) AAV engineering (scAAVengr) pipeline for rapid evaluation of transgene expression, which allows direct, head-to-head comparison of multiple vectors across all cell types, in parallel, in the same animals.7 We first validated the scAAVengr pipeline in NHP retina.

Although the NHP retina is highly similar to human retina, a direct correlation between the tropism of AAVs in NHP and human retina is yet unproven. It is therefore important to validate new AAV capsids in human tissue prior to clinical application.8–11 In this study, we applied the scAAVengr pipeline to human retinal explants in order to quantify the tropism of 18 wild-type and recently engineered AAV capsids with single-cell resolution. Simultaneously, we compared the expression profile of human ex vivo cultured retina with fresh retina. Our study further advances the understanding of the retinal ex vivo culture model and provides detailed information.

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AAV incubation was performed 1 day after culturing with 10 plants from the left and right eyes were cultured in separate wells. A total of 4.94 every second day with the complete replacement of the medium. A AAV library applied to the surface of each explant and was repeated canine-derived DE variants K91, K912, K916, and K947; NHP-derived DE variants NHP9, NHP26, and SCH/NHP2615; and 7m8, a variant created through DE in mouse retina6 (Table 1). Each variant was packaged into an AAV capsid. Individual AAV variants containing unique barcodes were pooled and the representation of AAVs in the library was quantified by deep sequencing. Little GFP expression was observed around blood vessels or in the fovea when it was located within the center of the explant. On day 8 of AAV incubation (day 9 of the explant culture), the highest infected region (edges) of each explant was dissected and collected. Each explant (total of 6 explants representing 3 regions from left and right eyes) was processed individually for single-cell RNA-seq (scRNA-seq). The single-cell suspension of macular, mid-peripheral, and peripheral retinal explants from the same eye were then combined, and GFP-positive cells were enriched by fluorescence-activated cell sorting (FACS; 2 FACS samples representing all 3 regions from left and right eyes) and processed for scRNA-seq. The remainder of the explant was fixed for imaging. Retinal explant cross sections showed that the basic structure of the retina was retained, with the presence of the retinal ganglion cell layer and inner nuclear and outer nuclear layers apparent in imaging. GFP was expressed across the retinal layers, with the strongest expression in photoreceptors in the outer retina (Figures 2E and 2F).

Figure 1. Illustration of the scAAVengr pipeline

(A) Generation of the AAV library. A CAG-GFP construct fused with a unique barcode was packaged into an AAV capsid. Individual AAV variants containing unique barcodes were pooled and the representation of AAVs in the library was quantified by deep sequencing. (B) AAV infection in the retina and transgene expression. Successfully infecting AAVs enter the nucleus and drive the expression of capsid-specific barcoded GFP mRNA in the infected cells. (C) The AAV-infected retina is processed for single-cell RNA-sequencing. Retinal tissue is dissociated, and the transcriptomes of single cells are sequenced. UMAP plots are used to visualize clusters of retinal cells, and the cell types of clusters are identified based on the expression of cell-type-specific marker genes. (D) The barcoded GFP transcripts are quantified, allowing for the tropism and infectivity of AAV variants to be evaluated across cell types with single-cell resolution.

about the performance of leading clinically relevant AAV serotypes in human retinal explants.

RESULTS

Construction of the AAV library
In order to provide quantitative information about the tropism and efficiency of promising AAV variants in human retinal tissue, we screened an 18-member AAV library on human retinal explants using the scAAVengr pipeline (Figure 1). The AAV variants in the library included naturally occurring serotypes AAV1, AAV2, AAV5, AAV8, AAV9, and AAVrh10; tyrosine- and threonine-mutated AAVs AAV2-4YF, AAV2-4YFTV, AAV8-2YF, and AAV9-2YF12–14; canine-derived DE variants K91, K912, K916, and K94; NHP-derived DE variants NHP9, NHP26, and SCH/NHP2615; and 7m8, a variant created through DE in mouse retina6 (Table 1). Each variant was packaged with a GFP transgene fused to a unique 25-bp barcode. The library was then created by pooling together the GFP-barcoded AAV variants, and the representation of each variant in the library was determined by deep sequencing (see methods).

Tropism of AAV variants in human retinal explants
The macular, mid-peripheral, and peripheral regions from the temporal quadrant of the retina of human donor #1 were dissected and cultured ex vivo with the photoreceptor side facing down on a transmembrane placed in a culture well (Figures 2A and 2B). Retinal explants from the left and right eyes were cultured in separate wells. AAV incubation was performed 1 day after culturing with 10 μL AAV library applied to the surface of each explant and was repeated every second day with the complete replacement of the medium. A total of 4.94 × 1011 vgs of AAV library in a volume of ~120 μL was added to each culture well over the course of 8 days of AAV incubation.

GFP expression appeared on the edge of the retinal explant at day 2 of AAV incubation, with an additional area of increased expression toward the center of the retinal explant. This expression pattern was the same for retinal explants from all anatomical locations, including macular, mid-peripheral, and peripheral retina, regardless of the presence of anatomical structures such as major blood vessels or the fovea (Figures 2C, 2D, and S1). Little GFP expression was observed around blood vessels or in the fovea when it was located within the center of the explant. On day 8 of AAV incubation (day 9 of the explant culture), the highest infected region (edges) of each explant was dissected and collected. Each explant (total of 6 explants representing 3 regions from left and right eyes) was processed individually for single-cell RNA-sequencing (scRNA-seq). The single-cell suspension of macular, mid-peripheral, and peripheral retinal explants from the same eye were then combined, and GFP-positive cells were enriched by fluorescence-activated cell sorting (FACS; 2 FACS samples representing all 3 regions from left and right eyes) and processed for scRNA-seq. The remainder of the explant was fixed for imaging. Retinal explant cross sections showed that the basic structure of the retina was retained, with the presence of the retinal ganglion cell layer and inner nuclear and outer nuclear layers apparent in imaging. GFP was expressed across the retinal layers, with the strongest expression in photoreceptors in the outer retina (Figures 2E and 2F).

scRNA-seq quantification of AAV efficiency across cell types in human retinal explants
Single-cell cDNA libraries created from macular, mid-peripheral, and peripheral retina regions and FACS cells from all retinal regions combined were deep sequenced, and transcripts were aligned and quantified. Data from the same anatomical location in the left and right eyes were combined, cells were clustered in a low-dimensional space, and cell type labels were identified for each cluster based on the most significant differentially expressed retinal cell-type marker genes. The numbers of cells analyzed after filtering were as follows: macula, 4,860; mid-periphery, 2,480; periphery, 3,131; and FACS, 3,162. Barcoded GFP transcripts originating from the various AAVs were then quantified and mapped to the identified cell types using the associated single-cell barcodes (Figure 3, and refer to methods for details).
Clusters of major retinal cell types including rods, cones, bipolar cells, ganglion cells, amacrine cells, horizontal cells, and glial cells were identified in the retinal explant samples (Figure 3). This is in agreement with scRNA-seq data performed with fresh retina from donor #2 (Figures S2A–S2C) and as previously reported. Similar quality-control statistics were observed between retinal explant samples and fresh retina samples from donor #2, with slightly lower unique molecular identifier (UMI) counts observed in the retinal explant samples (Figure S3). However, the cone population was only identified in the macular retinal explant samples, with the number of cones being lower than expected. The number of cones identified did not correlate with histological imaging. Cones were labeled with peanut agglutinin (PNA) but were swollen and significantly shortened in the retinal explants compared with freshly fixed retina from the same eye (Figures 4A–4D). This was especially apparent at the edge of the explants, where tissue was collected for scRNA-seq (Figures 4E–4G). Uniform manifold approximation and projection (UMAP) plots revealed a population of cells that did not express the cell-specific markers used for cell-type identification but had high expression of GFP transcripts (Figure 3, undefined cluster). This cluster maps closely to the rod and cone population and was enriched in FACS samples, which were largely populated with photoreceptors (Figure 3D). We thus hypothesized that this may be a group of photoreceptors with an altered transcriptome profile. We quantified expression of general photoreceptor gene markers as well as rod- and cone-specific gene markers (Figure S4; Table S2). Interestingly, as in rods and cones in the fresh retina samples, this undefined cluster had high expression of pan-photoreceptor markers, including Recoverin (RCVRN), guanylate cyclase activator 1A (GUCA1A), and VOPP1 WW domain binding protein (VOPP1) (Figures 3, S2, and S4). We observed no cone marker expression in this undefined cluster and only partial expression for some of the rod markers (PDE6A, GNGT1). In addition, in this undefined cluster, we found high expression of suppressor of cytokine signaling 3 (SOCS3), a well-known negative regulator of cytokine signaling that is involved in many cellular processes, including inflammation and cell death. In fresh retina samples, in contrast, the expression of SOCS3 is limited to the microglia (Figure S2). SOCS3 has been considered an indicator of retinal stress and has been shown to be upregulated during retinal degeneration. This evidence strongly suggests that degenerating photoreceptors make up this undefined cluster, hereinafter referred to as the SOCS3+ photoreceptor (SOCS3+ PR) population.

To further elucidate the SOCS3+ PR population, we integrated and clustered cells from all regions together (Figure S5). Interestingly, the SOCS3+ PR-labeled cells separate into two major clusters at this level. One of these populations clusters more closely with the small population of healthy cones and consists mostly of cells from the macula (57%). Cone densities vary according to eccentricity and are most concentrated in the macular retina, with a decrease in density toward the periphery of the retina. This population lacks expression of any cone-specific gene markers. Although it has minimal expression of the rod-specific gene markers, this can likely be attributed to low levels of overall ambient expression (background noise) from the large rod population, a common finding in retinal scRNA-seq data. The other population clusters closely with the rod population, consists of relatively equal amounts of cells from all regions (26.5% macula, 18.8% mid-periphery, 25.8% periphery, 28.8% FACS sample), and has higher expression levels of rod-specific markers. Thus, we hypothesize that each of these SOCS3+ PR populations corresponds to a cone- and rod-degenerating population, respectively.

The performance of individual AAV serotypes was then evaluated across cell types using three metrics: the absolute number of cells infected by each serotype (Figure 5A), the percentage of each cell type infected by each serotype (Figure 5B), and the level of transgene expression mediated by each serotype in the infected cells (Figure 5C). Each of these metrics was corrected by the AAV dilution factors determined by deep sequencing of the library. Heatmaps of these metrics revealed that AAV variants engineered through DE outperformed naturally occurring AAVs as well as tyrosine- and threonine-mutated AAVs across cell types and in explants from all three anatomical locations. PRs were the most efficiently infected cell type in all three regions, with the highest number of cells infected, the highest percentage of cells infected, and the highest mean transcripts of infected cells. This is in agreement with GFP expression observed in retinal cross sections (Figures 2E and 2F). Also, the majority of cells identified in FACS samples, where the GFP-positive cells were enriched, were SOCS3+ PR (Figure 3D).

Next, in order to rank the best-performing pan-retinal serotypes by cell type, variants were plotted by the mean transcripts per cell in infected cells versus the percentage of cells infected for each AAV serotype (Figure 6). Across retinal cell types and regions, of the canine-derived variants, K91 outperformed other engineered serotypes. Of the primate-derived variants, SCH NHP9/26 outperformed other variants. Among all AAVs, canine-derived variant K91 showed the highest infectivity at all retinal regions, closely followed by canine-derived variants K912 and mouse-derived variant 7m8. While the naturally occurring serotypes are overall less efficient, AAV5 was the most efficient of the naturally occurring serotypes tested in PRs (Figures 5 and 6).

We have previously shown that multiple AAVs can infect a single cell. In order to understand the dynamics of AAV infection in human retinal explants, UpSet plots with the number of cells infected by a particular combination of AAVs (the intersection size) and the

| Table 1. AAV variants in the library |
|-------------------------------------|
| AAV variants | Serotypes |
| Naturally occurring | AAV1, AAV2, AAV5, AAV8, AAV9, AAVrh10 |
| Tyrosine- and threonine-mutated | AAV2-4YF, AAV2-4YFTV, AAV9-2YF |
| DE* in canine retina | K91, K912, K916, K94 |
| DE in primate retina | NHP9, NHP26, SCH/NHP26 |
| DE in mouse retina | 7m8 |

*DE, directed evolution.
number of cells infected by a particular serotype (the set size) were generated (Figure 7). As many as 14 serotypes simultaneously infected a single cell. As revealed by the intersection size, more cells were infected by a single high-performing serotype (K91, K912, and 7m8; Figure 7, black-dots section) or by the combinations of two (Figure 7, red-lines section) or three (Figure 7, yellow-lines section) serotypes. Based on the set size, without AAV dilution factor correction, K912 infected the greatest number of cells, followed by K91 and 7m8.

Validation of top-performing AAV serotypes
In order to validate the top-performing AAV serotypes and to compare AAV performance ex vivo and in vivo, we tested individual AAV variants in NHP and human retinal explants. K91 was the best-performing serotype in human retinal explants, but this variant has previously been shown to underperform compared with other variants in vivo via intravitreal injection in NHP retina (Figures S6 and S7). K912 was the second-best-performing serotype in human retinal explants and the top-performing serotype among the set tested in vivo in NHP retina (Figures S6 and S7). We therefore packaged the CAG-GFP constructs into K91 and K912 and titer matched them for testing in NHP retinal explants. An additional experiment was performed to control for the orientation of the explants in the culture dish, as the medium containing AAVs contacts the side of explants attached to the transmembrane but does not submerge the explants. One set of retinal explants was cultured with the PR side attached to the transmembrane (PR down), and another set of explants was cultured with the RGC side attached to the transmembrane and the PR side facing upwards (PR up). The PR-up retinas were infected with either K91-CAG-GFP or K912-CAG-GFP, and the PR-down explants were infected with K912-CAG-GFP. A total of \( 2.45 \times 10^{10} \) vgs of AAV in a volume of 120 μL were applied to each group, as previously described, during 8 days of AAV incubation (each group consists of 3 explants, which were collected from central, mid-peripheral, and peripheral retina).

On day 2 of AAV infection, GFP expression was first observed on the edge of rhesus retinal explants in the K91 PR-down, K912 PR-down, and K912 PR-up groups. The GFP expression increased in intensity and area with time and plateaued in intensity by day 8 of AAV infection (Figures 8A–8C). The retinal explants were then fixed and sectioned. In the PR-down groups, both K91 and K912 drove efficient GFP expression across retinal layers, with the highest expression observed in the outer retina, in accordance with data from scAAVengr analysis in the human retinal explants (Figures 8D and 8E). Interestingly, the K912-infected PR-up explants also showed GFP expression mainly in the outer retina (Figure 8F). The tropism of K912 in rhesus retinal explant was similar in both culture orientations, suggesting that the orientation of the explant in culture does not influence tropism and that K912 has high affinity for PRs.
K912, which outperformed other variants both in vivo in NHP retina and ex vivo in human retinal explants, was further validated in retinal explants from another human donor (donor #3). 7m8, which infects human explants efficiently and is currently in use in multiple clinical trials and naturally occurring AAV2, the parental serotype of 7m8 and K912, also the first and only clinically approved serotype, were tested in parallel. The ex vivo culture of central, mid-peripheral, and peripheral retina and AAV application were performed as previously described. A total volume of 120-μL K912-CAG-GFP (1.26 × 10^{10} vgs), 7m8-CAG-GFP (5.58 × 10^{11} vgs), and AAV2-CAG-GFP (1.33 × 10^{10} vgs) were applied during 8 days of AAV incubation. K912- and 7m8-infected explants started to show GFP expression on the edge of the explants at day 2, while AAV2 (in a similar dose as K912)-infected retina showed GFP expression at day 4 (Figures 7G–7I). In retinal cross sections, the strongest GFP expression was observed in the outer retina for all three serotypes. K912 and 7m8 drove significantly stronger GFP expression than AAV2 (Figures 8J–8O).

**DISCUSSION**

Ex vivo retinal culture is the only available transduction model of mature human retinal tissue, and it has been widely used in retinal research, bridging the gap between in vitro cell/organoid culture and in vivo experimental animals. In this study, we applied the scAA-Vengr pipeline to human retinal explants, allowing for head-to-head evaluation of the tropism of 18 AAV capsid variants with single-cell resolution. These experiments provide quantitative information about the leading serotypes currently in development for clinical
In our forming panretinal virus, but K91 did not outperform parental gene expression in human and rhesus retinal explants, which allowed by K912 and 7m8. All three of these serotypes mediated efficient expression in NHP retinal explants, indicating that the tropism of AAVs in retinal explant culture is more comparable to subretinal injection than intravitreal injection. We therefore compared the expression pattern of AAVs in retinal explants with expression patterns from intravitreal and subretinal injections. In vivo, following intravitreal injection, the highest transgene expression is observed in the foveola, in a perifoveal ring of retinal ganglion cells, and in the peripheral retina in punctate areas near blood vessels. In retinal explants, no regional differences of expression patterns were observed, and we did not observe higher expression of GFP in the foveola and around blood vessels. This lack of AAV transduction in the foveola and around blood vessels may also be due to differences in the accessibility or presence of AAV receptors in these regions, and further histological studies will be required to better understand this differing expression pattern.

In our experiments, both the inner and outer retina of the explants have direct access to AAVs. The AAV suspension was added onto the RGC side of the explants and then quickly diffused into the medium, which is in contact with the PR side of the explants. As revealed by histological analysis, and with the scAAVengr workflow, the strongest GFP expression in retinal explants was observed in PRs. This expression was driven by a ubiquitous CAG promoter and was observed following administration of the AAV library and the individually tested serotypes. The same expression pattern in retinal explants has previously been reported by other groups.10,28

To better understand if the high AAV infectivity in PRs is a result of easier accessibility due to the orientation of the ex vivo culture system, we cultured rhesus retinal explants with either PRs facing down to the transmembrane or facing up and infected them with K912-CAG-GFP, a high-performing serotype, in both ex vivo human retina and in vivo NHP retina. The highest expression levels of GFP were observed in the outer retina in both culture orientations. This indicates that the tropism of AAVs in retinal explant culture is more comparable to subretinal injections than intravitreal injections in vivo regardless of culture orientation. Therefore, while retinal explants provide a valuable system in which to confirm AAV infectivity for human retinal cells, the usefulness of ex vivo culture may be limited for the prediction of tropism via intravitreal injection in vivo.

The lower efficiency of K91 in vivo via intravitreal injection may be due to lower ability to bypass the anatomical barriers that exist in vivo but are not intact in the ex vivo culture system. Retinal tissue in ex vivo culture maintains its normal gross morphological structure, as well as heterogeneous cell populations. However, anatomical structures including the vitreous and inner limiting membrane, barriers that restrict AAV diffusion in vivo, are not intact, especially on the edge of the explants. In explants from all anatomical locations including macular, mid-peripheral, and peripheral retina, AAVs showed higher expression levels near the edge of the explants than in the center of the explants, likely due to this lack of anatomical barriers at their edges allowing for easier accessibility of AAVs.

In a previous scAAVengr experiment, performed using intravitreal injections in NHP retina in vivo, K912 was identified as a top-performing panretinal virus, but K91 did not outperform parental serotypes. In our ex vivo human retinal culture, in contrast, K91 had better performance than K912, as quantified by the scAAVengr pipeline. To rule out the possibility that the different performance of K91 in human ex vivo tissue is due to cross-species differences, K91-CAG-GFP was packaged separately and used to infect NHP retinal explants. Transduction with K91-CAG-GFP drove high levels of transgene expression in NHP retinal explants, indicating that the improved performance of K91 is a result of the ex vivo culture system rather than any species specificity.

In retinal explants, no regional differences of expression patterns were observed, and we did not observe higher expression of GFP in the foveola and around blood vessels. This lack of AAV transduction in the foveola and around blood vessels may also be due to differences in the accessibility or presence of AAV receptors in these regions, and further histological studies will be required to better understand this differing expression pattern.

Figure 4. Unhealthy cones in retinal explants
(A) Freshly fixed human retina. (B) Human retinal explant at day 9 of culturing. (C) Freshly fixed rhesus retina. (D) Rhesus retinal explant at day 9 of culturing. (A–C) are all from the mid-peripheral retina. ONL, outer nuclear layer; INL, inner nuclear layer; RGC, retinal ganglion cell layer. (E–G) A flatmounted human retinal explant at day 9 of culturing. (F) Unhealthier cones (rounded, swollen) are observed at the edge. (G) The morphology of cones is better preserved (elongated) at the center.

Subretinal injections in vivo result in strong transgene expression in PRs under the injection bleb. Several AAV serotypes mediate high levels of transgene expression in PRs following subretinal injection in vivo. AAV5, for example, although it has low efficiency via intravitreal injection, drives fast-onset and efficient expression in PRs when administered into the subretinal space. As revealed by histological analysis, and with the scAAVengr workflow, the strongest GFP expression in retinal explants was observed in PRs. This expression was driven by a ubiquitous CAG promoter and was observed following administration of the AAV library and the individually tested serotypes. The same expression pattern in retinal explants has previously been reported by other groups.10,28
Additionally, both the scAAVengr pipeline and individual validation showed that K91 has high affinity for PRs in human and NHP retinal explants, and it may be of interest to further evaluate the potential of K91 for subretinal delivery in vivo.

Although human retinal explant culture is an important preclinical model, a significant drawback of this model is the limited survival window. As seen through histological analysis, all retinal cell types degenerate over time. However, it is not well understood how retinal ex vivo culture affects the expression profiles of retinal cells compared with fresh retina. Here, for the first time, we evaluated the condition of the human retinal explant culture at the single-cell level. Under the conditions tested here, 9 days after culturing, the major retinal cell types expressed similar retinal cell-type marker genes as the fresh retina except for a portion of the PR population, which showed an altered expression profile with the loss of cone- and rod-specific genes and upregulation of SOCS3, a potential indicator of retinal stress.

While SOCS3 is only expressed in microglia in fresh retina, upregulation of this gene was observed in rods and cones in the retinal explant. Previous studies have also noted that PRs are the fastest deteriorating cell types in ex vivo culture at the histological level and that the degeneration pattern is comparable to retinal injury and degenerative diseases. The fact that AAVs efficiently infect degenerating PRs in human retinal explants suggests that they may be promising for the treatment of retinal degeneration in patients.

In conclusion, we quantified the performance of 18 AAV serotypes in human retina ex vivo culture and determined that K91, K912, and 7m8 are the most efficient serotypes. Traditionally, the efficiency of AAVs has been evaluated by the expression level of transgenes such as GFP via histological analysis, a method that lacks quantitative accuracy. Using the scAAVengr workflow, we were able to simultaneously compare the efficiency of multiple AAVs in the same retinal explants and in the same cells. These results provide detailed

Figure 5. Quantitative comparison of AAV variant expression across retinal cell types
(A) Numbers of cells infected by each AAV variant. (B) Percentage of each cell type infected by each AAV variant. (C) Level of transgene expression driven by each serotype in the infected cells. Data are shown as mean transcripts per cell/100,000 transcripts. All data are corrected by the AAV dilution factor. Each plot was generated with 2 samples from the left and right eyes. PR, SOCS3+ photoreceptors; HC, horizontal cell; BP, bipolar cell; AC, amacrine cell; MiG, microglia; MuG, Müller glia; RGC, retinal ganglion cell.
Figure 6. Serotype performance across retinal regions

Scatterplots show the number of transcripts in infected cells per 100,000 transcripts versus the percentage of cells infected for each serotype. Best-performing variants that infect the highest number of cells and express the most GFP transcripts appear toward the upper right corner of each plot. All data are corrected by the AAV dilution factor. Each plot was generated with 2 samples from the left and right eyes. (A) Macula (B) Mid-periphery (C) Periphery (D) FACS-sorted cells. PR, photoreceptors; RGC, retinal ganglion cell.
information about leading AAV vectors for the preclinical development of gene-therapy approaches. Here, we have defined top-performing AAVs as those with the highest and widespread transduction across cell types. However, AAVs with cell-type specificity or a particular level of transgene expression may be required for other gene-therapy approaches, and the dataset provided here should enable the choice of the optimal AAV, out of the pool tested, for any particular therapeutic requirements. This study also provides valuable insight into the benefits and drawbacks of using retinal explant culture for the evaluation of AAV serotypes. Ex vivo culture is a useful model for confirming the infectivity of AAV in human retina, yet it may not be the best system with which to determine the tropism of AAV serotypes in vivo due to the discrepancies in infection patterns between in vivo and ex vivo models.

**Methods**

**AAV production and quantification**

A set of constructs containing unique 25-bp DNA barcodes after the stop codon of EGFP with a self-complementary CAG promoter and flanked by inverted terminal repeats (ITRs) was packaged separately using a triple transfection method using 293AAV cells (Cell Biolabs). AAVs were purified by iodixanol gradient ultracentrifugation, buffer exchanged, and concentrated with Amicon Ultra-15 Centrifugal Filter Units (#UCF8100) in DPBS. Each variant was pooled, and the titer of the virus was determined by quantitative PCR relative to a standard curve using ITR-binding primers or by using a QuickTiter AAV Quantitation Kit (Cell Biolabs). The relative titer of each variant in the pooled AAV library was confirmed by Illumina MiSeq sequencing (primer sequences are provided in Table S1).

**Postmortem human eyes**

All experiments were performed with approval and oversight from the University of Pittsburgh Committee for Oversight of Research and Clinical Training Involving Decedents. Eyes from postmortem donors were obtained through the Center for Organ Recovery & Education. Eyes was recovered from donors #1 (20-year-old male, within 3 h postmortem), #2 (58-year-old female, within 2 h postmortem), and #3 (45-year-old male, within 1 h postmortem) and then transferred to the lab within 30 min. All donors had no history of retinal disease.

**Rhesus macaque**

All procedures were performed in accordance with the Association for Research in Vision and Ophthalmology statement for the Use of Animals in Ophthalmic and Vision Research. All animal experiments were approved by the University of Pittsburgh Institutional Animal Care and Use Committee (IACUC). A 3-year-old male rhesus macaque was housed under standard 12-h light/12-h dark conditions.

For euthanasia, the animal was initially sedated with ketamine (15 mg/kg intramuscularly [IM]) and then ventilated and further anesthetized with isoflurane. The circulatory system was perfused through with 3–4 L ice-cold artificial cerebrospinal fluid (ACSF; 124 mM NaCl, 5 mM KCl, 2 mM MgSO4, 2 mM CaCl2, 23 mM NaHCO3, 3 mM NaH2PO4, 10 mM glucose; pH 7.4, osmolality 290–300 mOsm) oxygenated with 95% O2:5% CO2 at a rate of 120 RPM. Eyes were then enucleated and transferred on ice to the bench-top within 10 min.

**Retinal explant culture**

Eyes from postmortem donors/warehouse macaque was dissected, the anterior segment was removed, and eye cups were flat mounted. Around 5-mm wide macular (central), mid-peripheral, and peripheral retina was dissected, and retinal tissue was carefully separated from retinal pigment epithelium (RPE)/choroid/sclera and immediately transferred to cell culture inserts with 0.40-µm pore size (Thermo Fisher Scientific, #140640), with the PR side attached to the membrane (except for the PR-up group). Culture wells contained 3 retinal explants: macular (central), mid-peripheral, and peripheral retina. The retinal explants were cultured in Neurobasal Plus Medium (Gibco, A35829-01) supplemented with 2% B-27 (Gibco, A35828-01), 5 μg/mL Plasmocin prophylactic (InvivoGen, #ant-mpp), and penicillin-streptomycin (Genesee Scientific, PSL01-100 ML) at 37°C, 5% CO2. In order to keep the retinal explant attached to the transmembrane, a volume of 1.3 mL of medium was added into each culture well, which is a sufficient volume for wetting the transmembrane but not enough to submerge the explant. Fresh medium was replaced at the second day of incubation, and 10 μL AAV capsid library was applied dropwise onto the surface of each explant (30 μL per culture well). Every second day, medium was completely replaced, and AAV application was performed. A total of 120 μL of AAV suspension was applied per culture well during 8 days of AAV incubation. GFP expression in the retina was monitored using an ECHO Revolve fluorescence microscope. On the 8th day of AAV incubation, the 1-mm edges of each explant, which had the highest level of GFP expression, were collected and underwent single-cell dissociation, FACS, and scRNA-seq. Retinal explants used for histological analysis were fixed with 4% paraformaldehyde (PFA) for 2–4 h.

**Single-cell dissociation of retina tissue**

The 1-mm edge of each retinal explant was placed in Hibernate solution (BrainBits, HE500) after dissection, and was then dissociated with an enzymatic and mechanical method using a MACS Neural Tissue Dissociation Kit for postnatal neurons (Miltenyi Biotec, 130-094-802) according to manufacturer’s recommendations. The macular, mid-peripheral, and peripheral regions from the temporal quadrant retina of human donor #2 were dissociated, and the same anatomical

Figure 7. Cells infected by multiple AAV serotypes

The number of cells infected by a particular combination of AAVs (the intersection size) is shown in bar graphs across the top of each plot, and the number of cells infected by a particular serotype (the set size) is shown across the right-hand y axis. Dots and connecting vertical lines indicate the serotype and number of variants infecting a single cell. Lines are colored according to the number of AAV variants in the subset.
regions from the left and right eyes were combined and immediately dissociated using the same method. The retinal single cells were resuspended in D-PBS with 0.1% BSA and processed immediately for scRNA-seq or FACS.

**FACS**
The GFP-positive cells in the retinal explant cell suspensions were enriched using a MACS Tyto sorter (Miltenyi Biotec). GFP-positive cells were resuspended in D-PBS with 0.1% BSA and processed immediately for scRNA-seq.

**scRNA-seq and targeted gene enrichment**
Following the manufacturer’s protocol, using the Chromium Single Cell 3’ v.3 kit (10X Genomics), single retinal cells were partitioned into gel beads in emulsion (GEMs), and mRNA was reverse transcribed to cDNA and barcoded with UMIs. The GEM emulsions were then broken, and the cDNA was purified using DynaBeads. After PCR amplification, the cDNA was cleaned and further purified with SPRIselect reagent (Beckman Coulter, B23318). The final indexed library was constructed following fragmentation, end repair, A tailing, adaptor ligation, and sample index PCR steps. The libraries were sequenced on Illumina S1 flow cells at the UPMC Genome Center. GFP transcripts with unique AAV barcodes were further enriched using a custom Targeted Gene Expression kit (10X Genomics), and the resulting library was sequenced with an Illumina MiSeq reagent nano kit v.2 (300 cycles).

**scRNA-seq data processing and cell-type identification**
scRNA-seq data was aligned, and cells were demultiplexed using STARsolo (v.2.7). The GRCh38 reference genome (GCA_000001405.15) and its associated annotation file (NCBI RefSeq) were downloaded from UCSC and used for alignment. Empty droplets were removed using DropletUtils (v.1.4.3, lower.prop = 0.05). Doublets were removed using SCDS (v.1.0.0), with a hybrid score cutoff of 1.3. Gene expression was normalized using Scran (v.1.12.1), and imputation was achieved using ALRA (v.1.0) to improve the sparsity of the scRNA-seq datasets.

The analysis of scRNA-seq datasets and cell-type identification was performed with Scanpy (v.1.4.4.post1). Specifically, the top 50 principal components of the gene-expression matrix were calculated and used to compute the Euclidean distance between cells. Cells were embedded into a neighborhood graph for visualization using the UMAP algorithm, and Leiden clustering was performed on this graph to identify cell populations. Cell types were identified using a hypergeometric test using a list of known retinal cell-type marker genes compiled from multiple sources. p values were corrected for multiple hypothesis testing using the Bonferroni method, and the cell type for each cluster was chosen based on the most significant marker gene intersection p value (<0.05). Clusters that did not meet the significance threshold for cell-type identification were analyzed and annotated manually using the known gene markers.

All data from human donors, including cell-by-gene matrices, cell-by-AAV matrices, and fastq files, are available on GEO under GEO: GSE199840. Fastq files were deidentified using BAMboozle before being uploaded to the database.

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**Figure 8. Validation of individual AAV serotypes**
(A–C) GFP expression of K91- and K912-infected rhesus retinal explants 8 days post infection. Dotted line, edge of the retinal explants. (D–F) GFP expression in cross sections from the K91- and K912-infected rhesus retinal explants 8 days post infection. (G–I) Onset of GFP expression in the K912-, 7m8-, and AAV2-infected human retinal explants from donor #3. (J–O) GFP expression in the K912-, 7m8-, and AAV2-infected human retinal explant cross sections 8 days post infection. (J–O) were imaged with the same acquisition parameters. Each group has 3 explants from central, mid-peripheral, and peripheral retina. ONL, outer nuclear layer; INL, inner nuclear layer; RGC, retinal ganglion cell layer.
**Quantification of AAV barcodes**

AAV variants were identified using a unique 25-bp barcode at the end of the GFP gene for each variant. GFP was identified from these samples using two datasets: (1) whole-transcriptome scRNA-seq data and (2) 10x targeted enrichment against GFP and other marker genes. Salmon\(^\text{44}^\) (v.0.9.1) was used for GFP transcript quantification, and in-house scripts were used to identify the AAV barcode and map these reads back to their respective cells, as previously described.\(^7\)

**Immunohistochemistry**

The retinal explants remained on the transmembrane and were fixed in 4% PFA on ice for 2–4 h and dehydrated sequentially in 5%, 10%, 20%, and 40% sucrose solutions in PBS at room temperature (RT) (at least 30 min for each concentration). The transmembrane was cut around the retinal explant, and the retinal explants with the transmembrane were embedded in a 1:1 mixture of 40% sucrose in PBS:optimal cutting temperature (OCT) compound (Thermo Fisher Scientific) in liquid nitrogen for cryosection. Sections of 14 μm were collected on glass slides. Cryosections were rehydrated with PBS for 10 min at RT followed by PBS plus 0.1% Triton X-100 (PBST) for 30 min (retinal explant flatmounts were incubated in PBST directly at RT for 30 min) and then were blocked with 5% goat serum in PBST for 30 min. Primary antibody incubation was performed at RT for 1–2 h. Sections were washed with PBST, and then secondary antibody incubation was performed at RT for 1 h. For staining nuclei, Hoechst 33342 (Thermo Fisher Scientific) was diluted 1:5,000 in PBS and applied for 12 min. Retinal explant sections and flatmounts were imaged using an Olympus FV1200 confocal microscope. Antibodies used were rabbit anti-GFP (Thermo Fisher Scientific, A11122, 1:1,000), goat anti-Rabbit immunoglobulin G (IgG; H + L) Highly Cross-Adsorbed Secondary Antibody Alexa Fluor Plus 488 (Thermo Fisher Scientific, A32731, 1:1,000), and PNA (Thermo Fisher Scientific, L32460, 1:200).

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.omtm.2022.04.014.

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**AUTHOR CONTRIBUTIONS**

Z.X.: conceived, planned, and executed experiments, analyzed data, and wrote the manuscript; B.E.O., conceived, planned, and executed experiments, analyzed data, and wrote the manuscript; M.E.J., conceived, planned, and executed experiments, analyzed data, and wrote the manuscript; S.T., planned and executed experiments; W.R.S., supervised work, conceived, planned, and executed experiments, and wrote the manuscript; L.C.B., supervised work, conceived, planned, and executed experiments, analyzed data, and wrote the manuscript.

**DECLARATION OF INTERESTS**

W.R.S. is an inventor on a patent application on AAV screening methods. L.C.B. is an inventor on patent application on AAV capsid variants and AAV screening methods and is a founder of Newsight Therapeutics and Vegaent. All other authors declare no competing interests.

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