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Rod Outer Segment Development Influences AAV-Mediated Photoreceptor Transduction After Subretinal Injection

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Keywords
AAV, cones, gene therapy, photoreceptors, retina, rods

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Vectors based on the adeno-associated virus (AAV) are currently the preferred tools for delivering genes to photoreceptors (PR) in small and large animals. AAVs have been applied successfully in various models of PR dystrophies. However, unknown barriers still limit AAV's efficient application in several forms of severe PR degenerations due to insufficient transgene expression and/or treated cells at the time of injection. Optimizations of PR gene therapy strategies will likely benefit from the identification of the cellular factors that influence PR transduction. Interestingly, recent studies have shown that the AAV transduction profile of PRs differs significantly between neonatal and adult mouse retinas after subretinal injection. This phenomenon may provide clues to identify host factors that influence the efficiency of AAV-mediated PR transduction. This study demonstrates that rod outer segments are critical modulators of efficient AAV-mediated rod transduction. During retinal development, rod transduction correlated temporally and spatially with the differentiation order of PRs when vectors were introduced subretinally but not when introduced intravitreally. All subretinally injected vectors had an initial preference to transduce cones in the absence of formed rod outer segments and then displayed a preference for rods as the cells matured, independently of the expression cassette or AAV serotype. Consistent with this observation, altered development of rod outer segments was associated with a strong reduction of rod transduction and an increase in the percentage of transduced cones by 2- to 2.8-fold. A similar increase of cone transduction was observed in the adult retinal degeneration 1 (rd1) retina compared to wild-type mice. These results suggest that the loss of rod outer segments in diseased retinas could markedly affect gene transfer efficiency of AAV vectors by limiting the ability of AAVs to infect dying rods efficiently. This information could be exploited for the development of more efficient AAV-based PR gene delivery procedures.

Keywords: AAV, retina, photoreceptors, cones, rods, gene therapy

INTRODUCTION

Recombinant AAV serotype 2 (AAV2) is currently being evaluated in Phase 1/2 and 3 gene therapy clinical trials for treating diverse inherited retinal diseases involving the retinal pigment epithelium (RPE), such as Leber congenital Amaurosis 2, retinitis pigmentosa, and choroideremia. These studies demonstrate that subretinal delivery of AAV is safe and can provide rescue of vision-guided behavior in patients with advanced disease at least for several years. However, broader application of retinal gene therapy will require efficient transduction of other retinal cells, in particular photoreceptors (PR).

The ability of various AAV serotypes to transduce PRs has been evaluated in small and large animals. Among these, pseudotypes -2/5, -2/7, -2/8, -2/9, and -2/rh10 were found to have the highest transduction efficiencies for PRs after...
subretinal delivery.27,28 These vectors have been used to drive persistent transgene expression successfully, resulting in improvements of the retinal structure and/or function in a number of animal models of PR dystrophies, in particular in models of slow PR death.13 Yet, in models of severe PR degeneration, many PR therapies have not been very effective at rescuing PR over the long term. This is likely due to the difficulties in treating sufficient cells and/or expressing therapeautic levels of the transgene in a timely manner. In support of this idea, AAV8-based vectors have provided higher benefits than AAV5 in different disease models of primary PR dystrophies, such as in the retinal degeneration 10, the Rpgrip1 \(-/-\) (retinitis pigmentosa GTPase regulator interacting protein 1), the Gyer2e \(-/-\) (retinal guanylate cyclase 1), the Aipl1 \(-/-\) (aryl hydrocarbon interacting protein like 1), and the Rho \(-/-\) (rhodopsin) mice, consistent with the superior ability of AAV8 to transduce PRs more efficiently and thus delivering higher transgene levels when compared to AAV5 (for review, see Petit et al.13). Moreover, a recent study has shown that a therapeutic transgene can halt degeneration, regardless of the severity of PR loss if activated optimally in all PR cells.29 These results point to the importance of improving PR transduction efficiency to maximize clinical impact, and indicate that further development of retinal gene therapy will likely benefit from a better understanding of the potential host cell factors that restrict PR transduction in vivo. However, basic aspects of AAV–PR interactions are still unknown, and mechanisms affecting PR transduction after subretinal injection remain largely unexplored.20

Interestingly, recent reports have shown that after subretinal injection, the AAV transduction profile of PR cells changes dramatically between neonatal and adult mice.30–34 In particular, injection of AAV2/8 at postnatal day (PND) 0 results primarily in cone transduction (~100% of cones, almost no rods),32–34 whereas the vast majority of transduced cells in adults are rods.14,15,32 The current study took advantage of this shift in tropism to identify host factors critical for AAV-mediated PR transduction. Using 13 different AAV serotypes, cone and rod transduction were specifically assessed after AAV subretinal delivery in the developing retina and in two murine models of rod-cone dystrophies that display either an arrest of PR outer segment development or a rapid loss of rod cells. Irrespective of the serotype tested, it was found that all vectors preferentially transduced cones when delivered at PND1, whereas delivery at PND21 resulted in a decrease in the percentage of transduced cones and a dramatic increase in the percentage of transduced rods. Additional injections at PND5 and PND10 showed that rod transduction temporally and spatially correlates with the development of their inner and outer segments. Furthermore, in mutant mouse models that lack rod outer segments or rod PR cells, a preferential transduction of cones was observed. Together, these findings indicate that the access of AAV vectors to PR cells through their segments is a key factor for PR transduction upon subretinal delivery. Additionally, the data suggest that mature rods and cones may compete for AAV access. These observations begin to define a new paradigm for AAV–PR interactions upon subretinal delivery, which should impact the development of more efficient AAV-based PR gene delivery procedures by providing an understanding of host factors required for successful AAV transduction.

**MATERIAL AND METHODS**

**Plasmid construction, AAV vector production, and purification**

Recombinant AAV2/5-CMV-GFPd, AAV2/5-CMV-H2bGFP, AAV2/5-CMV-Cre, AAV2/5-pQCMV-H2bGFP, and AAV2/5-mCAR-H2bGFP were produced using the pAAV2-CMV-GFPd, pAAV2-CMV-H2bGFP, pAAV2-CMV-Cre, pAAV2-pQCMV-H2bGFP, or pAAV2-mCAR-H2bGFP plasmids, respectively. The pAAV2-CMV-GFPd plasmid expresses the destabilized GFP cDNA under the control of a human CMV enhancer/promoter, a human \(\beta\)-globin intron and an SV40 polyA signal, flanked by two AAV2 inverted terminal repeat sequences. The pAAV2-CMV-H2bGFP plasmid expresses a histone 2B fused GFP (nuclear GFP). It was constructed by replacing the GFPd cDNA with the H2bGFP sequence derived from the parental pQCMV-H2bGFP plasmid. The pAAV2-CMV-Cre plasmid expresses a Cre recombinase. It was constructed by replacing the GFPd cDNA with the Cre sequence derived from the parental pQCMV-H2bGFP plasmid. The pAAV2-mCAR-H2bGFP-I-Cre plasmid. The pAAV2-mCAR-H2bGFP plasmid was made by replacing the CMV promoter with the mouse cone arrestin (mCAR) promoter from the parental pQCMV-H2bGFP plasmid.35

scAAV1, -2, -3b, -4, -5, -6, -7, -8, -9, rh8, rh10, rh39, and rh43 expressing enhanced GFP (eGFP) under the control of the CB6 promoter were produced using the pAAVsc-CB6-eGFP plasmid, which bears a CB6 promoter, a rabbit globin polyA, and engineered ITRs for scAAV vector. The CB6 sequence was previously described36 and includes a CMV enhancer/beta-actin (CB) promoter with a CMV IE enhancer.
AAV2/5-CMV-GFPd, AAV2/5-CMV-H2bGFP, AAV2/5-CMV-Cre, AAV2/5-pQCMV-H2bGFP, and AAV2/5-mCAR-H2bGFP were produced in the Punzo laboratory by triple transfection of 293 cells according to previously reported methods and purified by two rounds of CsCl ultracentrifugation. scAAV vector production was carried out by the Vector Core of the Horae Gene Therapy Center of UMass Medical School (Worcester, MA). Viral vector titers were determined simultaneously for all batches by quantitative real-time PCR (qPCR) with primers directed toward SV40pA (forward: 5'-GGGGTTCCTTGTAGTTAATGA-3' and reverse: 5'-GGGTTCTTTTGTAGTTAATGA-3') or eGFP (forward: 5'-AGCAAGACCCCAACGAGA-3' and reverse: 5'-GGCCGGCCTACGAA-3') and expressed as vector genome per milliliter (vg/mL). The final vector titers were between 8.1×10^{12} and 5.1×10^{13} vg/mL (Table 1).

Animals

The CD1, Ai9 Cre reporter mice, and Pde6βrd1/rd1 mice were purchased from the Jackson Laboratory. The M-opsin-Cre mice (cone-specific Cre line) and the rhodopsin knockout (Rho-/-) mice were provided by Yun Z. Le (University of Oklahoma Health Sciences Center) and Janis Lem (Tufts University, Boston), respectively. Ai9+/MCre+ mice were generated by crossing MCre+ mice with the Ai9 Cre reporter mice. Pde6βrd1+/Ai9+/- mice were generated by crossing rd1 mice and Ai9 mice. Heterozygotes were mated to produce Pde6βrd1/rd1+/Ai9+/- mice.

All animals were maintained at UMass Medical School under a 12 hour/12 hour light/dark cycle with unrestricted access to food and water. Lighting conditions were kept constant in all cages, with illumination ranging between 10 and 15 lux. All experiments involving mice were conducted in compliance with the Association for Research in Vision and Ophthalmology statement for the use of animals in ophthalmic and vision research. All procedures were approved by the Institutional Animal Care and Use Committee of UMass Medical School.

Subretinal administration of rAAV vectors

The same experimenter (L.P.) performed all subretinal injections of AAV vectors. Subretinal injection of AAV vectors was performed using a trans-scleral approach through the posterior part of the sclera, as previously described, with the following modifications. Injections were performed using thin-wall beveled glass micropipettes (Clunbury Scientific LLC) without sclerotomy. In mice older than PND1, a small hole was made at the transition of the cornea and sclera with a 33-gauge needle before the injection in order to release intraocular pressure and allow for the formation of a vector bleb upon subretinal injection. This procedure allowed the vector bleb to occupy 40–60% of the retinal surface without injection-related damage. Fast green dye was added to the AAV preparations at a final concentration of 0.1% as a tracer to visualize the location of injection and thus ensure that AAV vectors were injected into the subretinal space. Mice received 0.5–0.75 l of vectors at PND1, 1–1.5 l of vectors at PND5–10, and 1.5–2.5 l of vectors at PND21.

Intravitreal administration of rAAV vectors

Intravitreal injections were performed in PND1 mice, as previously described, using the same glass micropipettes (Clunbury Scientific LLC) introduced through the cornea-scleral margin. Fast green dye was added to AAV preparations at a final concentration of 0.1% as a tracer to visualize the location of injection and thus ensure that AAV vectors were injected into the intravitreal cavity. Mice received 0.75–1 l of vectors.

Quantification of cone and rod transduction

In all cases, the analysis was performed 3 weeks post AAV injection. The total number of retinas analyzed in each experimental group is indicated in Table 1. The efficiency of retinal transduction was first assessed under an inverted fluorescent microscope, and only retinas that displayed native GFP fluorescence in >30% of the total retinal surface were processed for further analysis. All images were acquired with the Leica DM5500 fluorescent microscope equipped with a motorized stage for tiling and z-stack image acquisition and deconvolution software for confocal-like image quality. Quantification of cone and rod transduction in selected retinas was performed using three independent methods. First, for each group of injected mice, a minimum of two retinas were entirely sectioned (20 μm/section) and collected serially on five slides, such that each slide contained a representation of the entire eye. A minimum of two slides were processed for immunohistochemistry using Cy3-labeled peanut agglutinin lectin (PNA; 1:500; Vector Laboratories) and rabbit anti-cone arrestin (1:500; EMD Millipore). Images were taken from 5–10 sections at 40× over the entire transduced area at different focal plans (z stacks) using epifluorescence microscopy (Supplementary Fig. S1; Supplementary Data are available online at www.liebertpub.com/hum). The acquired epifluorescent images were then deconvolved in order to
obtain confocal-like resolution by eliminating the out-of-focus signal that could be misinterpreted as colocalization (Supplementary Fig. S1a). Cone quantification was performed in one selected z stack for each image, based on the colocalization of the GFP signal with the cone arrestin marker (usually at the level of cell nuclei). Rod quantification was performed within the same z stack for each image by manually counting the number of total GFP+ PRs and the number of GFP+ cones per image, and by determining through an automated counting algorithm the total number of PR nuclei in the outer nuclear layer (ONL) of the same image using the Imaris software package (Supplementary Fig. S1b). Rod transduction was then calculated as an average per eye and per group by determining the ratio of the number of total GFP+ PRs – number of GFP+ cones to the total number of PR nuclei – number of cone cells per image.

Second, for groups of mice injected with AAV-dGFP or AAV-eGFP vectors, cone quantification was also performed on retinal flat mounts stained with the following lectins/antibodies: Cy3-labeled PNA (1:500; Vector Laboratories), rabbit anti-cone arrestin (1:500; EMD Millipore; AB15282), rabbit anti-ALM opsin (1:500; EMD Millipore; AB5405), and goat anti-S opsins (1:500; Santa Cruz Biotechnology; SC-14363). For each flat mount, four to six representative 100 μm² areas within the transduced area (center and periphery) were analyzed. The percentage of transduced cones was determined as an average per eye and per group by quantifying the number of GFP+ cones over the total number of cones in the same areas. This quantification was performed at the level of cone segments, where GFP fluorescence in cones was unambiguously surrounded by PNA staining (Supplementary Fig. S1c).

Third, AAV5-CMV-H2bGFP and AAV5-pQCMV-H2bGFP vectors were also evaluated in Ai9+ MCre mice, which express CRE-mediated td-Tomato (red) in the entire cone cell bodies (Supplementary Fig. S1d).

**Cell lines**

Two different Chinese hamster ovary (CHO) cell lines were used to confirm the specificity of the lectins used in this study. These include the parental cell line Pro5 and sialic-acid deficient cell line Lec2. Both cell lines were kindly provided by Miguel Sena-Esteves (UMASS Medical School). Cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum and 1% streptomycin and maintained at 37°C with 5% CO₂. Cells were seeded at 10⁵ cells/well in 12-well plates prior to treatment for 1 h at 37°C with DMEM alone or DMEM supplemented with 50 mIU/mL of neuraminidase type III from *Vibrio cholerae* (Sigma–Aldrich; N7785). Cells were then washed three times with DMEM, once with PBS-1× and fixed with 4% PFA for 10 min. Cells were subjected to staining for 1 h in PBS-1× at room temperature (RT) using the following lectins: Cy2- or Cy3-labeled PNA (1:500), Cy2-conjugated *Maackia amurensis* lectin (MAL1; 1:500), biotinylated *Erythrina cristagalli* lectin (ELC; 1:500), and Cy5-conjugated *wheat germ agglutinin* (WGA; 1:500) all from Vector Laboratories. After three washes with ice-cold PBS-1× to remove unbound lectins, ECL was visualized using Streptavidin-Cy5 (Molecular Probes). Cells were imaged using an inverted fluorescence microscope.

**Lectin analysis of the inter-photoreceptor matrix**

Analysis of the inter-photoreceptor matrix (IPM) was performed on retinal sections, flat mounts, and explants from Ai9+ MCre+ mice euthanized at PND1, PND5, PND10, PND14, and PND21, and on retinal flat mounts from *Rho−/−* mice euthanized at PND21. Three eyes were used for each experimental group. Retinal sections and retinal flat mounts were processed and stained, as previously described, for the cell culture experiments, with the following panel of lectins in PBS1X: PNA (1:500; Vector Laboratories), MAL1 (1:500; Vector Laboratories), ELC (1:500; Vector Laboratories), and WGA (1:500; Vector Laboratories). For retinal explants treated with neuraminidase, retinas were dissected in cold DMEM and incubated in either DMEM alone or DMEM with 50 mIU/mL of neuraminidase type III from *V. cholerae* (Sigma–Aldrich; N7785) for 1 h at 37°C. Retinal explants were then washed three times with cold DMEM, once with cold PBS-1×, and fixed in 4% PFA for 30 min. Retinal explants were subjected to lectin staining for 1 h in PBS-1× at RT. After three washes with ice-cold PBS-1× to remove unbound lectins, retinal explants were flat mounted and analyzed by fluorescence microscopy (Leica DM5500).

**AAV-binding assay and qPCR**

AAV-binding assay was performed on three biological samples and repeated two to four times with retinal explants from PND1 and PND21 CD1 mice. Retinas were dissected in cold DMEM and incubated in either DMEM with or without neuraminidase for 1 h at 37°C. Retinal explants were then washed three times with cold DMEM, and pre-chilled at 4°C for 1 h in 500 μL of DMEM. AAV vectors were added at 1 × 10⁹ vg/well, and explants
were incubated at 4°C for 1 h. Retinal explants were then washed three times with cold DMEM and once with cold PBS-1× to remove unbound AAV particles. Total DNA was extracted using the DNaseasy Blood and Tissue Kit (Qiagen). AAV genome copies were quantified by real-time PCR with the aforementioned primers for the SV40pA or eGFP sequence. AAV vector genomes (vg) were normalized to the number of retinal cells, using primers complementary to the mouse beta actin DNA (forward: 5'-ACTGGGACGACATGGAGAAG-3'; reverse: 5'-GGGGTGTAGGTAGTCCTCAA-3'). Samples were run in triplicate. The results were expressed as mean AAV vg per genome DNA.

**Localization of AAV binding to retinal photoreceptors**

Localization of AAV particles was assessed after subretinal injection of PND1 and PND21 Ai9_MCre+ mice with scAAV8 or scAAV9 vectors. Eyes injected with PBS-1× were used as negative controls. One hour post injection, animals were euthanized and retinas processed for flat mount, as previously described.41 Retinas were incubated overnight at 4°C with polyclonal rabbit antibodies raised against intact capsids of AAV8 or AAV9 at 1:500 in blocking solution (kindly provided by G.G.). Sections were then incubated with a Cy2-conjugated secondary anti-rabbit antibody (Jackson ImmunoResearch; 1:500) for 2 h at RT. Results were analyzed by fluorescence microscopy (Leica DM5500).

**Statistical analysis**

All data are expressed as mean ± standard deviation (SD). All statistical comparisons use Student’s t-test, with p < 0.05 considered as statistically significant in all comparisons.

**RESULTS**

**Subretinal injection of AAV5 at PND1 predominantly targets cone photoreceptors**

Recent reports showed that subretinal injection of AAV8 into PND0 mice results primarily in cone transduction, whereas the vast majority of transduced cells after subretinal injection in adult mice are rods.32–34 To demonstrate further that PR postnatal development impacts the pattern of PR transgene expression, neonatal mice (PND1) were subretinally injected with 6×105–1.2×106 vg of single-stranded (ss) AAV2/5-CMV-dGFP (Table 1 and Supplementary Fig. S2a). AAV5 was chosen because this serotype (i) has been shown to significantly target PR cells in the adult mouse,14,15,18 cat,26 dog,23,42,43 pig,15,22 and nonhuman primate21; (ii) has been used successfully to restore retinal function and preserve retinal structure in various models of PR diseases13; and (iii) is currently planned in upcoming Phase I–II clinical trials for the treatment of PR diseases.13 Mice with successful subretinal injections were euthanized 3 weeks post injection (wpi), at PND21, after PR differentiation and maturation is completed (Fig. 1 and Supplementary Fig. S2). As previously described,30,33,34 subretinal injections at PND1 yielded GFP expression throughout the entire retinal surface (Fig. 1a and Supplementary Fig. S2a), presumably because the interactions between the RPE and the PR outer segments that could limit the spread of the vector have not yet formed at PND1. Robust transduction of RPE was observed across the retina (Supplementary Fig. S2b). High levels of GFP were also detected in the ONL, which contains rod and cone PR nuclei (Fig. 1c and Supplementary Fig. S2b), and in the corresponding PR segments (Fig. 1a and c and Supplementary Fig. S2b). Immunostaining with cone-specific markers (PNA and cone arrestin) revealed that the majority of the GFP+ cells were also positive for the cone-specific markers (Fig. 1a and c and Supplementary Fig. S2b). Hence, as predicted, the transduction of PRs was essentially relegated to cones when considering the percentage of GFP+ cells (cones: 69.1 ± 7.4%; rods: 2.7 ± 3.1%). L/M and S cones were equally transduced (Supplementary Fig. S3).

To rule out an influence of factors such as inhibition of CMV promoter activity or dGFP steady state level on the final transgene expression pattern, PR transduction was analyzed in mice injected at PND1 with five additional AAV5 vectors. These vectors carry different transgenes and/or ubiquitous or PR-specific promoters: AAV5-CMV-H2bGFP, AAV5-CMV-Cre, AAV5-mCAR-H2bGFP, AAV5-pQCMV-H2bGFP, as well as a self-complementary (sc) AAV5-CB6-eGFP (Table 1). Though some construction-dependency was noted, tropism was similar for all vectors examined (Fig. 1b and c and Supplementary Fig. S2b). Hence, the findings suggest that the absence of efficient rod transduction following subretinal administration at PND1 is not dependent on factors that alter vector genome expression, as well as second-strand synthesis in the transduced cells, but rather on differences in entry of AAV into cones and rods at the time of delivery. Consistent with that, AAV8 vectors carrying different promoters, such as AAV-hRK1-GFP, AAV-mCAR-GFP, or AAV-CMV-GFP, have also been seen to drive transgene expression predominantly in cones after subretinal injection in PND0 neonate mice.33,34 Moreover, efficient trans-
rod transduction after subretinal delivery

In mice, cones start to undergo their terminal mitosis to begin their subsequent differentiation before rods. Most rods in mouse are born over an extended developmental time period (E12 to PND5) with the peak of rod production around birth and their differentiation into mature rods continuing well into the postnatal period. To examine further how the PR transduction pattern correlates with rod PR development, GFP expres-
Figure 1. Photoreceptor transduction in the neonatal mouse retina by adeno-associated virus serotype 5 (AAV5) vectors. Mice injected on postnatal day 1 (PND1) were subretinally injected with similar doses of AAV2/5-CMV-dGFP, AAV2/5-CMV-H2bGFP, AAV2/5-CMV-CRE, AAV2/5-pQCMV-H2bGFP, AAV2/5-mCAR-H2bGFP, or scAAV2/5-CB6-eGFP vectors. (a) Representative retinal flat-mounts and (c, d) cryosections 3 weeks post injection (wpi) labeled with an antibody raised against anti-cone arrestin (CA) or with peanut agglutinin lectin (PNA; red) and counter stained with DAPI (blue). AAV-mediated GFP was detected by its native fluorescence (green). AAV-mediated CRE expression was detected in Ai9+ mice with the CRE-mediated expression of td-Tomato (white, native fluorescence). In retinal sections, retinal pigment epithelium (RPE) cells were artificially removed using Photoshop. Asterisks indicate GFP+ or CRE+ (td-Tomato+) cones, while arrowheads indicate GFP+ or CRE+ (td-Tomato+) rods, which are mainly found in the innermost rows of the outer nuclear layer (ONL). Quantitative analysis of the percentage of GFP+ or CRE+ photoreceptors in vector-exposed area is shown in (b). Error bars represent standard deviation (SD). Numbers in bars represent the number of retina analyzed. *p<0.05; **p<0.01; ***p<0.001 by Student’s t-test. (d) After injection of higher dose of AAV2/5-pQCMV-H2bGFP, a higher number of GFP+ rods is observed throughout the ONL (representative section). Scale bars = 25 μm. INL, inner nuclear layer; IS, inner segments; OS, outer segments; sc, self-complementary; ss, single-strand.
Figure 2. Efficiency of cone and rod photoreceptor transduction with AAV2/5 vectors at different times after birth. AAV2/5-CMV-dGFP, AAV2/5-CMV-H2bGFP, and scAAV2/5-CB6-eGFP were subretinally injected in mice at PND1, PND5, PND10, and PND21. In all cases, analysis was performed 3 wpi. (a) Representative retinal flat mounts (upper row) and cryosections (middle row) from mice injected with AAV2/5-CMV-dGFP and cryosections (bottom row) from mice injected with AAV2/5-CMV-H2bGFP and immunostained for PNA or CA (red) and counterstained with DAPI (blue; removed from 60% of panels to visualize red and green staining better). AAV-mediated GFP was detected by its native fluorescence (green). RPE cells were artificially removed using Photoshop. Dotted horizontal lines divide the ONL in three sublayers showing that early injections (PND1 and PND5) result in the majority of transduced rods (arrowheads) located in the most inner part of the ONL. Arrowheads indicate GFP+ rods (b). Quantification of cone, rod, and total photoreceptor transduction within the vector-exposed area after injection of AAV2/5-CMV-dGFP and AAV2/5-CB6-eGFP at the indicated ages. Results are shown as mean±SD. **p<0.01 and ***p<0.001 by Student's t-test. (c) Average distribution of GFP+ nuclei within the ONL layer after AAV injection at the indicated ages. As shown in (a), the ONL was divided into three sublayers: A, outermost part of the ONL; B, middle part of the ONL; C, innermost part of the ONL. The percentage of GFP+ nuclei/total nuclei (DAPI+) was calculated. Scale bars=25 μm.
The number of transduced rods increased, GFP+ S5a). After vector administration at PND5, while GFP at the time of injection, whereas the percentage of CMV- (Fig. 2a and b and Supplementary Fig. S4), AAV5- retina. Indeed, after injection of AAV5-CMV- rods that correlated with the differentiation of the revealed a progressive shift in tropism from cones to rods correlated with the differentiation of the retina. Indeed, after injection of AAV5-CMV-dGFP (Fig. 2a and b and Supplementary Fig. S4), AAV5-CMV-H2bGFP (Fig. 2a), or scAAV5-CB6-eGFP (Supplementary Fig. S5a and Fig. 2b), the percentage of GFP+ rods increased with the age of the mouse at the time of injection, whereas the percentage of GFP+ cones decreased. As a result, injections of the three vectors at PND1 and PND5 directed GFP expression predominantly in cones, while a preferential transduction of rods was observed for injections at PND10 and PND21 (Table 1).

Interestingly, the location of GFP+ rod nuclei in the ONL correlated with the timing of rod genesis (Fig. 2 and Supplementary Figs. S2b, S4, and S5a). After subretinal injection at PND1, the small percentage of transduced rod nuclei found was primarily located in the most inner row of the ONL where the first-born rods reside, as previously indicated by birth-dating experiments45–47 (Figs. 1 and 2a and c, and Supplementary Figs. S2b, S4, and S5a). After vector administration at PND5, while the number of transduced rods increased, GFP+ rods were still restricted in their location to the four or five most inner rows of the ONL (lower third of ONL; Fig. 2a and c and Supplementary Figs. S4 and S5a), consistent with the observation that newborn rods are stacked on the top of the earlier born ones.45 In contrast, injections after PND10 led to a wide distribution of GFP+ nuclei across the ONL (Fig. 2a and c, and Supplementary Figs. S4 and S5a). In line with this finding, the transduction pattern of rod and cone PRs followed the central-to-peripheral (data not shown) and ventral-to-dorsal gradient of PR differentiation48 when the vector was injected at PND3 (Supplementary Fig. S5b). Hence, the order of differentiation of PR cells is clearly correlated with the cells preferentially transduced after subretinal injection of AAV2/5.

Photoreceptor development affects rod transduction, irrespective of AAV serotypes

To determine whether PR development affects PR transduction by other AAV serotypes, neonatal mice were subretinally injected with a panel of 13 scAAV serotypes/variants, including AAV1, -2, -3b, -4, -5, -6, -7, -8, -9, -rh8, -rh10, -rh39, and -rh43, all expressing eGFP under the control of the CB6 promoter (Table 1, Figs. 3 and 4, and Supplementary Figs. S6 and S7). As controls, AAV5, -7, -8, -9, and -rh10 were subretinally injected into PND21 mice (Table 1 and Supplementary Fig. S6), as these vectors are known to exhibit excellent tropism for adult rods.14,15,18 Similar to the results observed after PND1 delivery of AAV5, PND1 injections of the all other AAVs resulted in preferential cone transduction when considering the percentage of GFP+ cells, with the exception of AAV4, which efficiently transduced RPE cells only (Table 1 and Fig. 3). On average, AAVrh10 exhibited the most efficient transduction of cones (although not statistically different from AAV5, -6, -7, -8, -9, -rh8, and -rh39), whereas AAV3b-mediated transgene expression was limited to sporadic RPE and PRs (Fig. 3b). Notably, no significant difference in the percentage of transduced rods was seen between all the serotypes (Fig. 3). When considering the percentage of GFP+ rods, all AAVs led to sparse rod transduction, with most of the GFP+ rods localized in the innermost part of the ONL (Fig. 4 and Supplementary Figs. S6 and S7). Additionally, the preferential transduction of rods observed after subretinal injection of AAV2/5 vectors at PND21 was reproduced after injection of AAV7 (Supplementary Fig. S6d and e), -8 (Supplementary Fig. S6a, d, and e), -9 (Supplementary Fig. S6b, d, and e), and -rh10 (Supplementary Fig. S6c–e). Thus, rod PR development is a critical host factor that influences AAV tropism after subretinal delivery independently of the serotype and thus probably independently of the different cell attachments factors utilized by each capsid.

Control of AAV transduction by photoreceptor development depends on the injection site

Because a higher number of GFP+ rods was observed after administration of higher doses of AAV5-CMV-H2bGFP (9 × 109 vg; Fig. 1d and Table 1) and AAV5-mCAR-H2bGFP (>1 × 1010 vg)41 at PND1, the overall absence of efficient rod transduction at PND1 may relate to the progressive loss of the episomal vector genomes in rods. AAV vectors integrate only at very low frequencies, and thus vector-mediated transgene expression is predicted to diminish over time under conditions of ongoing cellular proliferation. In transduced cones, the number of expressed vector genomes is likely to remain stable over time because cones are already post-mitotic by PND1–PND5.44 In contrast, although the vast majority of rod precursors are al-
Figure 3. Predominant transduction of cone photoreceptors after the injection of 12 different AAV serotypes/variants at PND1. (a) Representative images of retinal flat mounts at 3 wpi of scAAV1, -2, -3b, -4, -7, -8, -9, -rh8, -rh10, -rh39, or –rh43 vectors (all expressing eGFP under the control of the CB6 promoter), as indicated. Labeling with PNA (red) shows expression of eGFP (green) predominantly in cones (asterisks), except for AAV4. (b) Quantification performed on retinal flat mounts and/or cryosections of the percentage of GFP+ cones and GFP+ rods in the vector exposed area (note: actual transduction efficiency of scAAV4 was 0% for rods and cones). Results are expressed as mean±SD. Numbers in bars represent the number of retinas analyzed. *p<0.05; **p<0.01; ***p<0.001 by Student’s t-test. Scale bar = 25 μm.
ready post-mitotic by PND1\textsuperscript{44} and lineage studies using retroviral vectors have revealed that most rod progenitor cells that persist in the rodent retina after birth are terminally dividing,\textsuperscript{49–51} rod progenitors may dilute or lose AAV genomes, resulting in consistent decreased transgene expression in mature rods. Alternatively or concomitantly, the overall absence of efficient rod transduction at PND1 may relate to the progressive transcriptional silencing of AAV vector DNA due to developmental epigenetic changes. Indeed, histone modifications have been shown to be important contributors to gene regulation in developing rod PRs.\textsuperscript{52}

To investigate these two possibilities, AAV5, -6, -7, -8, and -9 were intravitreally injected in PND1 mice. If the factors that determine AAV tropism during PR development are related to AAV vector DNA, the final pattern of transgene expression should globally be similar between the two routes of administration. However, for all five serotypes, it was found that rods were the predominant PR cells transduced when the vectors were injected intravitreally (e.g., for AAV5: 4.2\textsuperscript{\pm}2.6\% of rods vs. 0.9\textsuperscript{\pm}0.3\% of cones near the injection site; \( n = 3 \); Fig. 4 and Supplementary Fig. S7). Notably, rod nuclei expressing detectable levels of GFP were evenly distributed across the ONL after intravitreal injection (Fig. 4 and Supplementary Fig. S7), indicating that lack of efficient AAV-mediated transduction of the late-born rods by subretinal injections at PND1 is not due to cell division or differential silencing of the vector ge-

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**Figure 4.** Comparative analysis of AAV vector transduction in the mouse retina after subretinal or intravitreal injection at PND1. PND1 mice injected either subretinally (upper row) or intravitreally (lower row) with scAAV5, -6, -7, -8, or -9 expressing eGFP under the control of the CB6 promoter, as indicated. Histological analysis was performed at 3 wpi on cryosections stained with CA (red) and counterstained with DAPI (blue). AAV-mediated GFP was detected by its native fluorescence (green). Subretinal injections result in a higher number of transduced cones (asterisks) with most of the transduced rods (arrowheads) located in the inner most part of the ONL (see also Figs. 1 and 2), while intravitreal injections result in scattered rods throughout the ONL and fewer transduced cones. Scale bars = 25 \( \mu \)m. GCL, ganglion cell layer.
nomenes. The data suggest that factors affecting the efficiency of AAV-mediated transduction of early and late-born rods are site specific and likely involve the access of AAV particles to the PR cells. Receptors necessary for binding/entry of AAVs while present in PRs may not yet be accessible on late-born cells by AAVs from the subretinal space.

**Lack of rod outer segments markedly alters PR tropism of AAV in the adult retina**

The timing of rod precursor birth (after E19) is tightly linked to the onset of PR segment formation and rhodopsin expression. To test whether the growth of rod segments and/or the associated matrix (Supplementary Fig. S8a and b) plays a role in the access of AAV particles to PRs, an attempt was made to visualize the distribution of AAV particles after subretinal injection at PND1 and PND21 using anti-AAV capsid antibodies. However, the large number of capsids present in the subretinal space limited the analysis (data not shown). Instead, qPCR was first employed to quantify viral binding on PND1 and PND21 retinas after modification of the rod IPM with neuraminidase (Supplementary Fig. S8). Ex vivo retinal explants were used to limit injection variability related to the surface of the retina exposed to the vector after subretinal delivery at PND1 versus PND21. We found that the rod-associated matrix could specifically interact with AAV particles in vitro (Supplementary Fig. S8d). However, these interactions were not conserved across AAV5, −7, −8, and −9, indicating that binding of AAV to the rod IPM does not govern the common increase in rod transduction observed during PR development after subretinal delivery.

Next, to determine if the development of a normal rod segment plays a role in the access of AAV particles to PRs, rod and cone transduction was quantified in rhodopsin knockout (Rho−/−) mice after subretinal injection of AAV2/5-CMV-dGFP, AAV2/5-CMV-H2bGFP, and AAV2/5-pQCMV-H2bGFP vectors at PND1 and PND21 (Table 1 and Fig. 5). The Rho−/− mouse is a well-characterized model of rod-cone dystrophy that does not develop rod outer segments. Importantly, this mouse model does not display any rod degeneration during the first three postnatal weeks. No alterations in the IPM and outer limiting membrane (OLM) were detected in Rho−/− mice at PND21 (Supplementary Fig. S9). In PND1-injected mice, the transduction profile was similar between Rho−/− and wild-type mice, with a preferential transduction of cone PRs (Fig. 5a and c). However, PND21 injections resulted in a fairly dramatic difference in the final expression patterns. On average, the three AAV vectors tested showed a similarly low transduction of rod PRs in Rho−/− retinas, as seen with PND1 injections in healthy retinas (Table 1 and Fig. 5a and c). Rod transduction was mainly observed at the site of injection near the needle tract (data not shown). GFP+ rods were located in the outermost part of the ONL (Fig. 5a, arrows) and expressed qualitatively less GFP than cone PRs (Fig. 5a and b). These observations suggest that the presence of a sick rod segment can influence the efficiency rod transduction upon subretinal delivery in the adult retina. Furthermore, they suggest a potential correlation between altered rod transduction and increased cone transduction.

To delineate further the influence of rods on the efficiency of cone transduction, the rd1 mouse model was utilized, in which the vast majority of rods are lost by PND21. However, in contrast to the Rho−/− retinas, the IPM is inevitably altered in this mouse model due to the loss of rods. AAV5 or AAV9 vectors were subretinally injected into PND21 rd1 mice and cone transduction was evaluated on retinal flat mounts at 3 wpi (Table 1 and Fig. 5d). All vectors resulted in dramatically enhanced cone transduction compared to control mice, considering the percentage of cells transduced (Table 1). The consistent finding in two mouse models of rod degeneration that cone transduction increases in the absence of efficient rod transduction or in the absence of rods suggest that rods may negatively affect cone transduction in the adult mouse retina.

**DISCUSSION**

Gene therapy targeting PRs holds great potential for the treatment of many forms of retinal diseases, such as retinitis pigmentosa, Leber congenital amaurosis, or age-related macular degeneration. A common hallmark of these diseases is the progressive dysfunction, degeneration, and death of PR cells. The first sign of these diseases—visual impairment—is generally associated with early alterations of the PRs functional structures, the inner and outer segments, while PR cell bodies are lost later in the process.

This study found that PR development and integrity play a major role in the efficacy of AAV-mediated transduction after subretinal injection. It confirmed that rod transduction increases dramatically during mouse postnatal retinal development, irrespective of the AAV serotype (Fig. 1 and Supplementary Fig. S6), and showed that rod transduction correlates with the differentiation order of PR cells (Fig. 2 and Supplementary Figs. S4 and S5). This effect was not due to differ-
Figure 5. Altered of AAV tropism in Rho−/− and adult rd1 mice. (a–c) PND1 and PND21 Rho−/− mice injected with similar doses of AAV2/5-CMV-H2bGFP, AAV2/5-CMV-dGFP, or AAV2/5-pQCMV-H2bGFP. Three weeks after injection, retinas were labeled with PNA or with an antibody raised against CA (red). AAV-mediated GFP was detected by its native fluorescence (green). Representative images after injection of (a) AAV2/5-CMV-H2bGFP and (b) AAV2/5-CMV-dGFP at age indicated. Asterisks indicate GFP+ cones and arrowheads indicate GFP+ rods. Quantitative analysis of the percentage of GFP+ cones (black bars) and rods (white bars) is shown in (c). Error bars represent SD. Numbers in bars represent the number of retina analyzed. **p < 0.01 and ***p < 0.001 by Student’s t-test. (d) Retinal flat mounts of rd1 mice injected at PND21 with similar doses of AAV2/5-CMV-dGFP, AAV2/5-CMV-H2bGFP, scAAV2/5-CB6-eGFP, or scAAV2/9-CB6-eGFP and stained for CA expression (red) 3 wpi. AAV-mediated GFP was detected by its native fluorescence (green). Percentage of transduced cones for each vector is indicated in corresponding panel as mean± SD. Scale bars = 25 μm.
ences in promoter activity or second-strand synthesis of the viral genome, which in principle could account for the enhanced rod transduction (Fig. 1 and Supplementary Fig. S2). Neither was this effect related to the gradual exit of rod progenitors from the cell cycle or to epigenetic/protein inhibition of the vector genomes during development, since both early and late-born cells were efficiently transduced at PND1 after intravitreal delivery of AAV vectors (Fig. 4). Thus, while serotype-independent (Figs. 3 and 4, and Supplementary Figs. S4–S6), the effects of PR development on AAV transduction were, however, site specific (Fig. 4).

It is possible that restriction to AAV transduction in immature rods occurs due to inefficient viral trafficking. However, intracellular AAV particle trafficking has been shown to be influenced by the viral capsid sequence, which in this case seems to be relevant with regards to the common shift in tropism between PND1 and PND21 subretinal injections. Alternatively, the enhanced transduction of rods during retinal development could be related to increased entry of AAVs into PRs. The OLM, in which the zonulae adherentes pore size has been estimated to be between 30 and 36 Å, may impede the diffusion of AAV capsids (250 Å) to the ONL, allowing efficient access (and thus entry) of AAV particles mainly to PRs that have started to develop inner segments that protrude through the OLM (i.e., cones and early-born rods). In support of this model, the timing of rod precursor birth (after E19) is known to be tightly linked to the onset of PR segment formation and rhodopsin expression. Concomitantly, the development of rod outer segment may directly or indirectly correlate with increased—nonspecific—entry of AAV vectors into PRs. Indeed, a shift in the distribution of GFP rod nuclei was observed when AAV5 vectors were delivered after PND10, which coincides with the elongation of rod outer segments for the majority of rods at this age (Fig. 2c). Moreover, rod transduction was profoundly inhibited in adult Rho−/− retinas (Fig. 5) that lack rod outer segment development but still display rod inner segments that protrude through the OLM. Of course, other factors, such as PR stress and degeneration, could affect the expression of AAV receptors, as well as inhibit the expression of the transgene in the diseased retina. However, there is more rational evidence from the observations to suggest that vector access to PR segments is one of the host factors governing PR transduction efficiency upon subretinal delivery, in particular since at PND21, Rho−/− retinas had no apparent loss of rod PRs or alterations of the IPM and OLM integrity (Supplementary Fig. S9).

Importantly, the idea that PR outer segments modulate rod transduction in the adult retina does not contradict previous reports that demonstrated unequivocal (but transient) structural and/or functional rescue of rods by AAV-mediated gene replacement therapy in two mouse models of retinitis pigmentosa that fail to develop rod outer segments; namely the Rho−/− and the Phrp2−/− mice. In the Rho−/− mice, therapeutic vectors were injected into neonatal mice and mice younger than PND5—two time points that precede rod outer segment growth in wild-type mice and where it has been shown that rod transduction is dependent on the viral load injected (Fig. 1d). In the Phrp2−/− mice, it has been notably shown that there was a significant difference in the number and quality of outer segment rescue in treated retinas. Depending on the age at which the mice were treated, fewer outer segments developed normally, with the best results obtained in younger animals, despite the relatively slow rate of PR death in that model. Moreover, particular differences in the infectivity of the vector preparations and the presumed vector titers may exist between all these studies, making it difficult to cross-compare them directly. The goal of this study was to determine host cell factors that influence PR transduction, and oversaturating the system with AAV vectors would have been counterproductive in this case. As mentioned, it has previously been shown in wild-type mice that the majority of immature rods in addition to inner nuclear layer cells can be transduced by subretinal injection at PND1 if a higher dose of vector is used (Fig. 1d). In this study, the rate of rod transduction after injection at PND21 was limited to 30–55%, while other groups have reported PR transduction rates of >90% within the vector exposed area. Thus, the overall lower transduction in this study can easily explain why in Rho−/− mice such a large difference with PND21 subretinal injections was seen. Another factor that complicates the comparison between experiments in the Rho−/− mice and the previously published results by Palfi et al. is that the present study used different Rho−/− mice. Strain background differences between the two strains and the way both knockout strains were generated could have exacerbated the effect of the lower transduction rate used to perform this study. In this regard, it is interesting to note that the group of Jean Bennett has also reported difficulties in transducing efficiently rods in the same Rho−/− model that was used in this study (Dejneka NS et al., ARVO Annual Meeting Abstract, 2002).
The present findings provide important information regarding the effects of subretinal gene therapy in animal models of PR dystrophies. First, the results indicate that the time of intervention affects the efficacy of AAV transduction if gene therapy is applied before the full development of the retina. However, in several mouse models of PR dystrophies, the degeneration of PRs is so fast that most gene therapy attempts have been conducted during the first week of life due to the relative slow onset of AAV-mediated transgene expression. For instance, in the nmf363 murine model of PDE6x-deficiency, no apparent PR degeneration is observed at PND12, but by PND14, 30% of PRs are already lost and only one row of PR nuclei remains in the ONL by PND38. In this model, injection of an AAV8(Y788F)-RHO-mPde6β at PND5 resulted in an initial loss of cells between 1 and 2 months of age, followed by a preservation of three to four rows of PR nuclei in the vector-exposed area, for at least 6 months. However, the overall rod functional rescue was too low to make a detectable difference by electroretinography. Notably, mid-stage intervention at PND21 achieved similar efficacy as PND5 treatment, despite loss of approximately half of PRs at the time of injection, indicating that the efficacy of PND5 treatment may have been limited by the efficient transduction of rods when compared to the PND21 intervention. As well, another study reported that subretinal injection of AAV2/5-smCBA-mPde6β vector (1 × 1010 genome copies) in rd10 mice at PND4 (before the onset of PR dystrophy) or PND21 (after the onset of PR dystrophy) resulted in similar (partial) therapeutic effects.

Second, the results indicate that if PR integrity determines the ability of AAV vectors to target the PRs, the spatio-temporal kinetics of retinal degeneration, the site of injection, and the health status of the retina at the time of treatment are also important factors affecting the overall efficacy of transduction. For instance, murine models of severe PR dystrophies show significant loss of PR segments at early stages of the disease. In these models, most preclinical studies have demonstrated improved PR survival only when gene therapy is applied before or at very early stages of retinal degeneration. Moreover, when gene therapy has been applied at later stages of the disease, the proportion of PRs that were not responding to the treatment (“silent” cells) increased within the vector-exposed area. It has been first suggested that there is a threshold of accumulated changes after which PR death is inevitable. However, recent data indicate that continued PR loss may instead reflect insufficient transduction efficiencies. An intriguing possibility is that AAV vectors might not efficiently access and infect subpopulations of deteriorated PRs. Thus, early alterations of PRs may be associated with a higher heterogeneity of transduction within the vector-exposed area (e.g., 40% of PRs expressing high levels of transgene and 60% of PRs expressing low levels of transgenes rather than 100% of PRs expressing medium levels of transgene). An important step in approaching this problem will be to understand the longitudinal changes that occur during phases of PR stress/degeneration at the level of both the PRs and AAV vector pharmacology. Future studies will also have to be designed to determine whether a temporal disruption of the OLM and higher doses of vectors may be able to overcome this hurdle. Finally, delineating the relationship between PR development, AAV trafficking, and cell receptor usage will be essential toward developing a complete understanding of PR transduction upon subretinal injection.

It did not escape the authors’ attention that the results support the notion that rods and cones may directly or indirectly compete for AAV access. Preclinical studies with several AAV serotypes have established that subretinal injection in adults often results in dominant transduction of rods within rod-rich retinas. In wild-type mice (cone:rod ratio of 1:30), while fairly high transduction of rods is observed after subretinal injection of AAV2/8, cone transduction is restricted to only 1–12% of cells. In pigs, AAV2/8 mediates 3.8- to 5- and 1.7-fold higher levels of PR transduction than AAV2/5 and AAV2/9, respectively. However, all serotypes transduce the same percentage of cones in the cone-enriched visual streak (cone:rod ratio of 1:3 to 1:5), with up to 9.2% of cones readily transduced. In dogs, AAV2/5 also preferentially targets rods, though significant cone transduction is observed in cone-enriched areas, with the use of high doses of vectors. Primate studies with seven different AAV serotypes reported similar findings with weak cone transduction in the rod-rich parafoveal region (1–10%), even in the presence of ample rod PR transduction, and with higher levels of cone transduction (20–50%) in the pure-cone fovea. Consistent with the notion that there is a developmental effect in which access by AAV to cone PRs becomes restricted, a profound decrease in cone transduction was observed in the developing mouse retina (Fig. 2). These observations appear to correlate with the increase in rod transduction (Fig. 2b). Moreover, cone transduction was significantly enhanced after subretinal injection of AAV vectors in PND21 Rho+/− (Fig. 5a–c) and rd1 mice (Fig. 5d), in which rod transduction was inhibited by the absence...
of a healthy rod segments or by the loss of the vast majority of rod cells at the time of treatment. Although comprehensive analysis will be required to elucidate the details of PR transduction upon subretinal injection, it should be noted that there is no evidence from the present observations to suggest that AAV particles use different receptors between rods and cones. Indeed, a similar shift in tropism was observed for all tested AAV serotypes (Fig. 3 and Supplementary Fig. S6). It is possible that the densely stacked rod outer segments form physical barriers between the site of delivery and the cones. Concomitantly, cone matrix sheaths located around cone inner and outer segments may form a barrier that selectively reduces access of AAV particles to the cones. Interestingly, a recent study in the feline retina has shown that unlike mice and nonhuman primates, cone PRs were more efficiently transduced than rods. The reasons for this difference in cellular tropism remain unknown, but the cone matrix sheath of the cat differs significantly from other mammalian species. In primates, only AAV9 was shown to target cones both centrally and peripherally efficiently at low doses when directly compared to five other AAV serotypes. It has been hypothesized that this property is due to the abundance of terminal galactose, the cell receptor for AAV9 on the cone PR matrix.

The notion that rods and cones compete for AAV transduction emphasizes the need of evaluating components of AAV vectors planned for humans in models that accurately depict physiological characteristics of the human retina (i.e., large animals and all-cone murine retinas). Indeed, while in humans macular cones will often be the primary treatment area, the ability of AAV vectors to transduce the cones and restore their function is often evaluated in murine models of PR dystrophies that do not display a cone-enriched area but a high rod:cone ratio. It also indicates that loss of rod PRs associated with many forms of retinopathies could be exploited to redirect AAVs commonly toward cones.

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AUTHOR DISCLOSURE

G.G. is a founder of Voyager Therapeutics, specialized in AAV-based gene therapy for the central nervous system, and holds equity in the company. G.G. is an inventor of patents with potential royalties licensed to Voyager Therapeutics and other biopharmaceutical companies. No competing financial interests exist for the remaining authors.

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