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Original Article



Mannose-coupled AAV2: A second-generation AAV vector for increased retinal gene therapy efficiency

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Inherited retinal diseases are a leading and untreatable cause of blindness and are therefore candidate diseases for gene therapy. Recombinant vectors derived from adeno-associated virus (rAAV) are currently the most promising vehicles for in vivo therapeutic gene delivery to the retina. However, there is a need for novel AAV-based vectors with greater efficacy for ophthalmic applications, as underscored by recent reports of dose-related inflammatory responses in clinical trials of rAAV-based ocular gene therapies. Improved therapeutic efficacy of vectors would allow for decreases in the dose delivered, with consequent reductions in inflammatory reactions. Here, we describe the development of new rAAV vectors using bioconjugation chemistry to modify the rAAV capsid, thereby improving the therapeutic index. Covalent coupling of a mannose ligand, via the formation of a thiourea bond, to the amino groups of the rAAV capsid significantly increases vector transduction efficiency of both rat and nonhuman primate retinas. These optimized rAAV vectors have important implications for the treatment of a wide range of retinal diseases.

INTRODUCTION

Recombinant viral vectors derived from adeno-associated viruses (rAAVs) constitute the most promising *in vivo* gene transfer platform for gene therapy of various human genetic diseases, including hemophilia, ^{1,2} muscular dystrophies, ³ and retinal blindness.⁴ Voretigene neparvovec (Luxturna), one of the six rAAV products currently on the market, is approved for the treatment of RPE65-related Leber congenital amaurosis, a congenital retinal blinding disease.^{4,5} The retina is an ideal candidate organ for gene-based therapies: it is readily accessible for surgical injection, can be imaged non-invasively in real time, responds well to local treatments, and its blood-retinal barrier limits unintentional spread of vectors to neighboring tissues and the general circulation.^{6,7} Moreover, the retina is a post-mitotic tissue in which gene transfer induced by non-integrating vectors can achieve long-term production of therapeutic protein. Finally, the rela-

tively immunoprivileged environment of the eye in general,⁸ and the retina in particular, may protect rAAV vectors from neutralization by pre-existing circulating anti-AAV antibodies.

However, clinical trials increasingly report adverse events in cohorts of patients locally injected with the highest rAAV doses.⁹ Inflammation and increased intraocular pressure were described in the high-dose cohort in a recent clinical trial of diabetic macular edema (NCT04418427) patients. With the development of more effective viral vectors for retinal gene delivery, therapeutic thresholds could be reached with lower vector doses, thereby reducing the likelihood of the aforementioned adverse events.

The most common strategy to optimize rAAV vector efficacy involves modification of the viral capsid via genetic engineering, using either rational design or directed evolution approaches. Rational design approaches seek to introduce targeted changes in the capsid based on prior knowledge of capsid structure and function.^{10,11} By contrast, directed evolution approaches consist of repeated cycles of random mutations or peptide insertions guided by desired biological traits rather than existing knowledge.^{12,13} Although these strategies are promising,¹⁴ manufacture of new modified rAAV vectors is

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challenging as it requires optimization of each stage of the production and purification processes, as well as complex characterization of the newly generated particles.

We have developed a chemical engineering approach to rAAV vector optimization based on the presence of specific basic amino acids on the AAV capsid. The use of a specific ligand could improve the targeting and transduction of specific cells but also modify the physico-chemical properties of the AAV, potentially increasing its efficacy. We previously described how functionalization of amino groups of rAAV2 capsids with a sugar bearing an isothiocyanate coupling function increases hepatocyte transduction in vitro with respect to non-modified vectors.¹⁵ A significant advantage of this strategy is that we use "natural recombinant AAV vectors," meaning that no genetic modification is required to produce the starting product. In the present study, to specifically target retinal cells, we synthesized a new ligand, an arylisothiocyanate mannose derivative (Figure S1). This sugar was chosen because it can modify the physico-chemical properties of molecules or biomacromolecules after chemical bioconjugation by increasing their hydrosolubility.¹⁶ We decided to use the AAV2 serotype based on our experience of its limited retinal diffusion, with transduction limited to the region of the bleb after subretinal injection.¹⁷ As a proof-of-concept mannose grafting on the capsid of AAV2 may increase the diffusion and also the number of transduced cells. We assessed the efficacy of these chemically modified vectors following subretinal injections in rats and nonhuman primates (NHPs). This new approach to the development of rAAV vectors has important implications for the rational design of safer and more effective AAV vectors for ophthalmic and other applications.

RESULTS

Synthesis of mannose ligands

We hypothesized that the development of rAAV particles carrying mannose carbohydrates would modify the physico-chemical properties of rAAV particles by increasing their solubilities and allow an increase of the transduced cells. Two sets of mannose ligands were designed and synthesized: one with a phenylisothiocyanate function capable of reacting with lysine residues bearing a primary amino group; and a second that lacked any reactive function and served as a control.

The chemical synthesis of the arylisothiocyanate (Ar-NCS; compound 5) is shown in Figures S1A. In the first step, the per-acetylated mannose compound 1 was glycosylated in the presence of BF₃-Et₂O and 2-[2-(2-chloroethoxy)ethoxy]-ethanol to obtain a significant yield of compound 2 in an exclusive α anomer configuration. The azide compound 3 was synthesized by nucleophilic substitution of the chloride of 2 in the presence of sodium azide and potassium iodide in dimethylformamide. This intermediate was reduced under hydrogen atmosphere with Pd/C in the presence of *p*-toluenesulfonic acid while deprotection of the hydroxyl groups was achieved using the basic resin IRN78 to obtain a quantitative yield of compound 4. Finally, the Ar-NCS compound 5 was obtained by reacting 4 with an excess of *p*-phenylenediisothiocyanate in dimethylformamide with 44% yield. Compound 7, which lacked any coupling function, was synthesized in two steps from commercial D-mannose pentaacetate, with an overall yield of 71%. This control was synthesized to demonstrate that only covalent coupling on rAAV capsid surface occurs with ligand 5 and to control for any passive adsorption of the mannose-based compounds to the capsid surface during the chemical coupling. Glycosylation of 1 in the presence of BF₃-Et₂O and 2-(2-ethoxyethoxy)-ethanol and cleavage of the acetyl groups resulted in the formation of the non-reactive 7 (Figures \$1B). All chemical compounds were characterized by NMR, HPLC, and mass spectrometry (Figures \$2–\$13).

Bioconjugation of mannose ligands to the AAV capsid

Bioconjugation of the mannose ligands 5 and 7 to the rAAV2 capsid surface was performed using an optimized version of a previously described protocol.¹⁵ Chemical coupling was achieved by nucleophilic addition of the amino group of the capsid proteins to the reactive isothiocyanate motif of ligand 5, yielding a thiourea linkage between the rAAV2 and the mannose ligand. Two different molar ratios of 5 (3×10^5 [3^E5] and 3×10^6 [3^E6]) were used per AAV particle to evaluate (1) the possibility of modulating the number of mannose ligands on the rAAV capsid surface and (2) the impact of this modulation on the therapeutic index of these new vectors. Compound 7 was used only at the highest molar ratio (3^E6) to demonstrate the absence of adsorption to the capsid surface (Figure 1A).

Western blot was first used to validate bioconjugation between rAAV2 and ligand 5. Capsid-specific polyclonal antibody detection confirmed the integrity of the VP capsid subunits after the chemical reaction (Figure 1B) for all conditions tested. Figure 1C shows the efficient grafting of ligand 5 onto the rAAV2 capsid using concanavalin A lectin (which recognizes mannose sugar), demonstrating the covalent coupling of 5 onto the three AAV capsid proteins (VP1, VP2, and VP3).¹ Conversely, rAAV2 capsid subunits incubated with ligand 7 could not be detected using concanavalin A lectin, confirming that the Ar-NCS function is essential for bioconjugation (Figure 1C). The amount of ligand 5 covalently attached to the three VPs clearly increased when the molar ratio was increased from 3^E5 to 3^E6 (Figure 1C), as reflected by the increase of the intensity of the bands. The purity and integrity of these mannosylated VPs was verified by silver staining, which showed that (1) VP1, VP2, and VP3 remained intact after the chemical coupling and (2) the molecular weight of each VP increased when the molar ratio was increased from $3^{E}5$ to $3^{E}6$ (Figure 1D).

Together these findings clearly demonstrate efficient bioconjugation of the mannose ligand **5** to the rAAV2 capsid via a thiourea covalent bond.

Aggregation of rAAVs is a well-described phenomenon.¹⁹ We therefore evaluated this parameter after bioconjugation. The size of rAAV2 and mannose-coupled rAAV2 particles was evaluated by dynamic light scattering (DLS) (Table S1). As shown in Table S1, the size of all particles ranged from 25 to 29 nm, indicating that the coupling of mannose ligands to the rAAV2 capsid did not promote aggregation.



Figure 1. First covalent coupling of 5 onto the capsid of AAV2 via primary amino groups

(A) $1.4^{E}12$ vg of AAV2-GFP vectors were added to a solution of compound **5** (3^E5 or 3^E6 eq) in TBS buffer (pH 9.3) and incubated for 4 h at RT. The same experimental procedure was followed with compound **7** (3^E6 eq) in TBS at pH 9.3 as a control. (B and C). 5^E8 vg of the samples were analyzed by western blot using a polyclonal antibody against the capsid to detect VP proteins (B) or using an FITC-concanavalin A lectin (C) to detect mannose sugar. Note that compound **7** is a negative control lacking the Ar-NCS reactive function. (D) 1^E10 vg of each condition were analyzed by silver nitrate staining. VP1, VP2, and VP3 are the three proteins constituting the AAV capsid. Capsid protein molecular weight is indicated at the right of the images according to a protein ladder.

tively). For evaluation of the first batch of mannose-coupled rAAV2, rats received injections of unmodified and mannose-coupled

To demonstrate the reproducibility of the bioconjugation process, AAV coupling was repeated using new batches of rAAV2 and ligand 5 ($3^{E}6$ eq). As shown in Figure S14 and Table S1, all findings (qPCR, western blot, silver staining, and DLS) were comparable with those obtained for the first batch, confirming the reproducibility of this technology.

As already described for rAAV2-GalNAc vectors, chemical modification of the capsid surface can impact on the infectivity of the modified particles.¹⁵ Indeed, chemical modification of lysine amino acids on rAAV2 capsid motifs may limit or even abrogate targeting of the recombinant virus to heparan sulfate, one of its common receptors. Consequently, the tropism of the chemically modified rAAV2 vectors could be altered, allowing transduction of cells that are usually nonpermissive to rAAV2, while reducing the transduction of cells that express heparan sulfate receptors on their surface. This hypothesis is supported by the relative transduction efficiency observed for all mannose-coupled rAAV2 vectors carrying a GFP reporter gene when evaluated in HEK293 and HeLa cells. As shown in Table S1, vector genomes (vg) per GFP-forming units (vg/GFU) values were in the same range for rAAV2, rAAV2 + 5 ($3^{E}5$ eq), and rAAV2 + 7 $(3^{E}6 \text{ eq})$, and about one log higher for rAAV2 + 5 ($3^{E}6 \text{ eq}$). Comparable results were obtained for the two coupling experiments.

In vivo evaluation in rats

To evaluate the *in vivo* efficacy of the mannose-coupled rAAV2 vectors for gene transfer in the retina, two distinct batches of mannose-coupled rAAV2 vectors were injected in two groups of rats (n = 18 and 15, respectively) (Table S2). In both groups, control rats received subretinal injection of 2.5 μ L vehicle (i.e., the vector formulation buffer). The experimental groups received 2.5 μ L of either rAAV2 (at a dose of 1^E12 vg/mL) or mannose-coupled rAAV2 (rAAV2 + 5 [3^E5] or rAAV2 + 5 [3^E6]) at doses of 1^E12 or 4^E11 vg/mL, respectively (total viral genomes injected per eye: 2.5^E9 and 1^E9 vg, respectively

rAAV2 vectors into the right and left eyes, respectively, and were monitored up to 4 weeks post-injection. For evaluation of the second batch of mannose-coupled rAAV2 vectors, rats received injections of vehicle, rAAV2, or mannose-coupled rAAV2 + 5 ($3^{E}6$) in both eyes (n = 5 animals per experimental condition). Eve fundoscopy was used for in vivo monitoring of GFP transgene expression post-injection at time points up to 6 weeks (Table S2). Retinal structure was analyzed by spectral domain-optical coherence tomography (SD-OCT) (Figures S15A). SD-OCT images of all eyes obtained before injection and immediately before sacrifice (i.e., 4 or 6 weeks post-injection) revealed preservation of the different retinal layers in all areas scanned. The thickness of the ONL is the same before and after injection in the injected area (Figures S16A). This finding is in agreement with a previous report in which no significant toxicity was observed for rAAV vector doses in the same range as used in the present study.^{20,21} Fundoscopic examination of retinas from eyes injected with unmodified rAAV2 (n = 28) or mannose-coupled rAAV2 vectors (rAAV2 + 5 $[3^{E}5]$, n = 9; rAAV2 + 5 $[3^{E}6]$, n = 19) revealed similar findings: GFP fluorescence was detected as early as 1 week post-injection, increased up to 6 weeks post-injection (the latest time point examined), and was limited to the retinal bleb induced by subretinal injection (Figures 2 and S17). A greater fluorescence intensity was quantified from D14 in retinas injected with rAAV2 + 5 $(3^{E}6)$ (n = 10 eyes) versus those injected with identical doses of unmodified rAAV2 (n = 10 eyes) (Figure 3). This difference in fluorescence intensity was not observed in retinas injected with rAAV2 + 5 $(3^{E}5)$ (n = 9, data not shown).

To determine which cell types were transduced after subretinal injection of rAAV2 (n = 18 eyes) or mannose-coupled rAAV2 vectors (rAAV2 + 5 [$3^{E}5$], n = 9 eyes; AAV2 + 5 [$3^{E}6$], n = 9 eyes), vertical sections were labeled with a fluorescent GFP antibody and overlaid with Mayer's hemalum staining (Figure 4). Vertical sections were selected at random to determine the relative tropism of modified



Figure 2. Ocular fundus in rats (first coupling)

Kinetics of GFP expression after subretinal delivery of AAV2, AAV2 + **5** ($3^{E}5$), or AAV2 + **5** ($3^{E}6$), determined using *in vivo* imaging. Images show representative acquisitions obtained with each vector type. The green color becomes saturated on AAV2 + **5** ($3^{E}6$) as the expression is very high, and the sensitivity of image acquisition on the retinophotograph is fixed for all retinal photographs from 1 week post-injection (wpi).

versus unmodified vectors, although it should be noted that this approach does not provide a quantitative measure of transduction (Figure 4).

We observed that the same cell types were transduced by both modified and unmodified rAAV2 vectors: in both groups GFP was detected in the retinal pigment epithelium (RPE) and outer nuclear layer (ONL). Weaker transduction of the inner retinal layer and ganglion cells was observed.

These findings indicate that chemical modification of rAAV2 with ligand 5 ($3^{E}6$) did not modify rAAV2 tropism in the retina, but did result in higher GFP expression, which could have a beneficial impact on vector efficacy, thereby allowing for a reduction in vector dose. To investigate the basis of this apparent difference in transduction efficacy, we performed molecular biology analyses of retinas injected with unmodified and chemically modified rAAV2. We quantified vector copy number by qPCR (Figures S18A and S18B) and GFP mRNA transcripts by RT-qPCR (Figures S19A and S19B) 4 weeks post-injection in neuroretinas and the RPE from rats injected with vehicle (n = 3), unmodified rAAV2 (n = 5), or modified rAAV2 + 5 ($3^{E}6$) (n = 5) vector.

In the neuroretina, vector genome copy numbers were significantly lower in rats injected with rAAV2 + $5(3^{E}6)$ than those injected with unmodified rAAV2 (mean 0.02 vector genome copy number per diploid genome [vg/dg] and 0.10 vg/dg, respectively; p = 0.0079). In the RPE, vector genome copy numbers were comparable regardless of the injection administered. However, in both the neuroretina and the RPE, GFP mRNA transcript levels were significantly (approximately 10-fold) higher in samples from rats injected with $rAAV2 + 5(3^{E}6)$ than in those injected with unmodified rAAV2 (mean relative quantity [RQ] = 0.268 and 0.039, respectively in the neuroretina, p = 0.0079; and mean RQ = 0.244 and 0.017, respectively in the RPE, p = 0.0397). Next, to evaluate the "activity" of one vector genome copy, we calculated the ratio of GFP mRNA levels (RQ) to vector copy number (vg/dg) in the neuroretina (Figure 5A) and RPE (Figure 5B). In the neuroretina we observed significant differences between rats injected with unmodified rAAV2 vectors and those injected with the modified rAAV2 + 5 (3^E6) vector: a single copy of the rAAV2 + 5 ($3^{E}6$) vector expressed up to 35 times more GFP mRNA than a single copy of the rAAV2 vector (Figure 5A, mean RQ/vg for rAAV2 = 0.4, and mean RQ/vg for rAAV2 + 5 $[3^{E}6]$ = 13.6; p = 0.0079). Note that this graph shows the ratio of transcript to vector genome. Hence, in a given retinal cell just one vector genome from the mannose-coupled vector can yield on average 17 transgene transcripts while it requires two vector genomes from the unmodified vector to have the probability of producing just one transcript. Similarly, in RPE samples GFP mRNA levels corresponding to a single copy of rAAV2 + 5 $(3^{E}6)$ vector were higher than those of the rAAV2 vector (mean RQ/vg for rAAV2 = 0.7, and mean RQ/vg for $rAAV2 + 5 [3^{E}6] = 4.8$), although this difference was not significant, likely due to the high variability of RQ/vg ratios in rats injected with the mannose-coupled rAAV2 vector (Figure 5B).

Taken together, these data showed that, compared with unmodified rAAV2, mannose-coupled rAAV2 vector exhibits similar cellular tropism but far superior transgene expression efficiency in the rat neuroretina.

In vivo evaluation in NHPs

Compared with rodents, the NHP retina has a thicker vitreous, a thicker inner limiting membrane, and a specialized fovea area with high visual acuity.²² We tested the efficacy of modified mannose-coupled rAAV2 vectors for retinal gene transfer in NHPs.

Three (n = 3) NHPs received subretinal injections of vehicle, rAAV2, or chemically modified mannose-coupled rAAV2 (rAAV2 + 5 [$3^{E}6$]) in the left eye, between the RPE and photoreceptor layer (Table S2). Subretinal injections were performed using a transvitreal approach, following vitrectomy with air exchange, following the same surgical technique used for the administration of Luxturna²³ in patients with RPE65-mediated inherited retinal dystrophy. Surgery was well tolerated by all animals. A vector concentration of $4^{E}11$ vg/mL was used, allowing for injection of a total viral load per eye of between $6^{E}10$ and $7^{E}10$, a dose conventionally used in preclinical and clinical settings.^{23–25} Animals were followed up for 4 weeks after injection. Retinal structure was analyzed by SD-OCT before injection and



immediately before euthanasia (Figures S15B). The slight decrease of thickness of the ONL is observed in three retinas (vehicle or vector injected) and therefore seem to be related to surgical procedure. However, this result should be confirmed with a larger number of primates (Figures S16B).

Expression of the GFP reporter was monitored by fundus fluorescence imaging before injection and at 1, 2, 3, and 4 weeks post-injection (Figure 6). As expected, no GFP fluorescence was observed in the eye injected with vehicle only. Fundoscopic examination of NHPs that received rAAV2 and rAAV2 + 5 ($3^{E}6$) vector revealed GFP fluorescence as early as 1 week after vector delivery, after which a continuous increase in fluorescence was observed up to at least 4 weeks post-injection. For the same vector dose, we observed greater fluorescence intensity in the retina injected with the mannose-coupled rAAV2 vector and a saturation of the signal as early as 15 days post-injection (Figure 7) supporting our observations in the rat model (Figure 3). To evaluate the cellular tropism of rAAV2 and chemically modified rAAV2, retinas were flat mounted and stained to label both cell nuclei (Draq5, in purple) and cone photoreceptors (LM and S opsin, in red) (Figure 8). The intensity of staining in retinas treated with chemically modified rAAV2 was far greater than for the nonmodified rAAV2, such that the direct quantitative comparison of the two retinas cannot unfortunately be made without saturation of the signals from the AAV2 + 5 $(3^{E}6)$ -injected retinas. Nonetheless, in these two primates, there were no evident differences in the cell types transduced by the mannose-coupled rAAV2 vectors compared with unmodified rAAV2 vectors. Transduced cells included retinal ganglion cells, cells of the outer plexiform layer and the ONL, and photoreceptor inner segments. Almost no transduced cells were detected in the inner nuclear layer or in photoreceptor outer segments. This tropism was consistently observed across the entire transduced area of retina. In agreement with eye fundoscopy images, immunohistochemistry revealed higher GFP expression in retinal flat mounts from NHP injected with chemically modified mannose-coupled rAAV2 (Figure 8). However, the proportion of the transduced area was similar in the retina injected with mannose-coupled rAAV2 vectors (57%) and unmodified rAAV2 vectors (60%) (Figure S20). Given the increased retinal transduction efficiency of chemically modified rAAV2, we used qPCR to compare the biodistribution in distant tissues (liver, spleen, occipital cortex, and optic chiasm) of the mannose-

Figure 3. Fluorescence intensity of GFP expression

-O-Vehicule -⊡-AAV2 -■- AAV2 +5 (3E6) GFP intensities on the fundus imaging measured at different time points post-injection in rats injected with vehicle (n = 6 eyes), rAAV2 (n = 10 eyes), and rAAV2 + 5 (3^E6) (n = 10 eyes). Results are expressed as mean ± SEM.

coupled rAAV2 vector versus unmodified rAAV2 following subretinal delivery (Table S3). Low numbers of vector genome copies (0.002 vg/dg; lower limit of quantification [LLOQ] of our qPCR assay, 0.001 vg/dg) were detected only in the occipital cortex of the animals injected with

rAAV2 + 5 ($3^{E}6$). No vector genome copies were detected in any of the other samples from NHPs injected with rAAV2 or rAAV2 + 5 ($3^{E}6$) vector. These results confirm previous reports describing an absence of unmodified rAAV biodistribution in the liver after subretinal delivery.^{26,27} Importantly, these data indicate that no extraocular distribution of the chemically modified mannose-coupled rAAV2 occurs after subretinal delivery in NHPs.

DISCUSSION

In this study, we have demonstrated a novel biosynthetic approach to generate a new generation of rAAV chemically coupled enhanced vectors for retinal gene therapy. Ocular gene transfer is a unique system for the treatment of inherited retinal disorders. However, and despite a first Food and Drug Administration-approved rAAV gene therapy product for a retinal genetic disease in 2017, no other ocular gene therapy has been yet authorized despite the dramatic increase of the number of clinical trials over the last decade.^{28,29} This may be due in part to the efficacy and safety drawbacks recently reported in preclinical and clinical studies. Emerging evidence suggests that ocular gene transfer is more inflammatory than believed previously.^{27,30–33} Despite a proven efficacy of voretigene neparvovec, the post-marketing trial PERCEIVE launched in 2019 followed up 103 treated patients and showed among them that 35% experienced at least one ocular adverse event including chorioretinal atrophic change mainly at the injection site, foveal degeneration, eye inflammation, or increased intraocular pressure.34 The inflammation described in dose-escalation studies reveals the higher titer cohort to be most at risk.³⁵ This dose dependence supports the need of new-generation rAAV vectors with an improved balance between efficacy and safety.

The novel rAAV2 vectors described here demonstrate altered transduction capacity both *in vitro* and *in vivo* through modification of the density of mannose ligands on the vector capsid surface. We show that subretinal delivery of mannose-coupled rAAV2 vectors into rat and NHP eyes in our study is comparable with that previously described in rodent models injected subretinally with native rAAV2 vectors.^{36,37} Importantly, we observed significantly higher transgene expression in rats injected with the chemically modified mannosecoupled rAAV2 vector. The results obtained in primates are in agreement with those found in rats but would need to be confirmed in a larger number of animals and with lower doses to allow an accurate



Figure 4. Vector distribution in the retina of rats

Four weeks after subretinal injection with AAV2, AAV2 + 5 ($3^{E}5$), or AAV2 + 5 ($3^{E}6$), from first coupling. Representative images of paraffin-embedded retina sections stained with anti-GFP antibody and anti-mouse Alexa 555-conjugated antibody (red labeling), and counterstained with hematoxylin. Scale bars, 30 μ m. GC, ganglion cells; INL, inner nuclear layer; ONL, outer nuclear layer; IS/OS, inner and outer segments; RPE, retinal pigment epithelium.

quantification of transduced GFP+ retinal cells on retinal sections. In NHP, we administered $6^{E}10$ to $7^{E}10$ total viral genomes per eye ($2^{E}8$ to $5^{E}11$ vg/eye being injected in patients) and this resulted in almost 100% of photoreceptors transduced in the subretinal bleb as shown by our retina flat mounts. It would have interesting to inject 1 or 2 log less of the vector to see a difference in the number of transduced cells between the two conditions (modified and unmodified vectors). Such quantitative data should be provided in future preclinical studies.

We chose the AAV2 capsid to demonstrate proof-of-principle in the serotype that is most well characterized across a range of tissues. However, other capsid types, both natural and engineered, could in principle be similarly chemically modified to increase transgene payload with the goal of reaching a therapeutic threshold with less virus.

Capsid molecular engineering through amino acid mutagenesis or directed evolution has previously resulted in AAV capsid variants with enhanced properties after ocular gene transfer such as AAV2TYF,³⁸ D4-R100,³⁹ or 7m8.¹² The two latter enable retinal gene transfer via intravitreal injection and are being used in the clinic.⁴⁰ Despite its surgical advantage, rAAV delivery via the intravitreal route was shown to be more immunogenic because of a reduced immune privilege and a higher systemic dissemination.^{27,41} With their enhanced transduction properties, our chemically engineered new rAAV allows the reduction of vector doses in the retina, which could reduce the risk of dose-related rAAV toxicity in treated patients, a phenomenon increasingly reported in clinical trials.

Another advantage over molecular engineering is the use of native rAAV as the starting product, which can rely on the current and already approved manufacturing platforms. A dramatic dose reduction should also decrease manufacturing needs and related costs, Moreover, altering the density of mannose ligands on the rAAV vector surface could have an impact on rAAV immunogenicity itself, with a reduced induction of fewer anti-capsid neutralizing antibodies because capsid antigenic determinants are hidden by the engrafted ligands. This was demonstrated previously in the liver for GalNac ligands on the surface of AAV vectors and will have to be investigated for other types of capsid-conjugated chemical compounds.¹⁵ Ligand density on the surface of the capsid is, however, one critical parameter to determine because it may influence vector transduction efficiency. Our *in vitro* data show indeed a higher vg/GFU ratio using rAAV2

coupled to ligand 5 at 3^E6 ligand payload may reflect reduced infectivity of these modified vectors: the higher ligand 5 payload may block the capsid motifs that are most efficiently recognized by the heparan sulfate receptors, thereby decreasing transduction efficiency in cells that bear these receptors. This is confirmed by the fact that this decrease was not observed for the rAAV2 + 5 ($3^{E}5$ eq) vector, perhaps because the lower coating density resulted in reduced blockade of capsid motifs. In line with in vitro data, we have observed comparable findings in vivo. The theoretical number of AAV2 capsid surface sites accessible for chemical ligand conjugation was estimated in a previous study.¹⁵ According to the AAV2 sequence, there are 18 lysine residues (out of 34) located in VP3, 8 of them being exposed on the capsid external surface. Considering that there are 60 VP3 proteins per particle, this gives a total of 480 amino groups potentially accessible for chemical coupling on the AAV2 capsid. As discussed above, we have shown that we can modulate the number of ligands on the capsid surface and subsequently the efficacy of the variant in the target cells. However, it is difficult at this stage to precisely determine the number of molecules grafted per VP and their precise sites on the capsid surface sites. Thus, quality control methods are still necessary to allow such structural characterization and the translation of such new rAAV products to the clinics.

In conclusion, altogether, our *in vitro* data and *in vivo* results in two animal species showed that, compared with unmodified rAAV2, mannose-coupled rAAV2 vector exhibits similar cellular tropism but far superior transgene expression efficiency in the retina. These modified mannose-coupled AAV2 vectors represent a new generation of biotherapeutic products for more efficient gene transfer in the retina application. They may offer an alternative to the current dose-related safety issues reported in patients and their efficacy is currently addressed in subsequent studies using preclinical models of retinal genetic disorders. In addition to the retina, this chemical modification of rAAVs could be applied to other serotypes relevant for ocular diseases and more largely to the central nervous system to further improve local delivery of rAAV vectors in these organs thanks to an enhanced efficacy at reduced vector dosing.

MATERIALS AND METHODS

Materials

All chemical reagents were purchased from Acros Organics or Sigma Aldrich and were used without further purification. Rabbit polyclonal

anti-AAV capsid proteins antibody (cat. no. 61084) was obtained from PROGEN Biotechnik. Anti-fluorescein-alkaline phosphatase (AP) Fab fragment antibody (cat. no. 11426338910) for the detection of fluorescein-labeled lectin was obtained from Sigma-Aldrich. FITCconcanavalin A lectin was purchased from Vector Laboratories. Reactions requiring anhydrous conditions were performed under nitrogen atmosphere. All chemically synthesized compounds were characterized by ¹H (400.133 or 300.135 MHz), ¹³C (125.773 or 75.480 MHz) NMR spectroscopy (Bruker Avance 300 Ultra Shield or Bruker Avance III 400 spectrometer). Chemical shifts are reported in parts per million (ppm); coupling constants are reported in units of Hz. The following abbreviations were used: s, singlet; d, doublet; t, triplet; q, quartet; quin, quintet; br, broad singlet. When needed, ¹³C heteronuclear HMQC and HMBC were establish structures. High-resolution mass spectra (HRMS) were recorded with a used to unambiguously Thermo Fisher hybrid LTQ-orbitrap spectrometer (ESI⁺) and a Bruker Autoflex III SmartBeam spectrometer (MALDI). HPLC analyses were performed on an HPLC autopurification system (Waters) equipped with a BGM 2545 binary pump, a 2767 Sample Manager, and a UV-vis diode array detector and evaporative light scattering detector in series (Waters 996 PDA and Waters 2424 ESLD). Chromatographic separation was performed on an Atlantis T3 (Waters; 4.6×150 mm, 5 μ). The mobile phase consisted of water (solvent A) and acetonitrile (solvent B). Gradient mix: 0.0 min [95% A; 5% B], 20.0 min [77% A; 23% B]. Flowrate: 1 mL/min.

All chemically synthesized products were purified by flash chromatography (Grace Reveleris Flash Chromatography System) equipped with UV and DLS detectors.

Synthesis

15 min. $F_3B \cdot OEt_2$ (1.90 mL, 15.384 mmol, 3 eq) was then added dropwise and continually stirred for 8 h at room temperature (RT). The mixture was diluted with DCM and washed with an aqueous saturated solution of NaHCO₃. The crude mannosyl glycoside **2** was used in the next step without further purification (2.78 g).

Compound 3

To a solution of the crude 2 in dry DMF (30 mL), NaN₃ (1.33 g, 20.512 mmol, 4 eq) was added and the mixture was stirred at 70°C under N2 atmosphere for 16 h. DMF was evaporated under vacuum conditions, and the residue was dissolved in AcOEt and washed with water and brine. The crude product was purified by column chromatography (SiO₂, Cy/AcOEt, 70/30 as eluent) to produce the azide derivative 3 (1.81 g, 3.581 mmol, 70% in two steps), a colorless oil. ¹H NMR (CDCl₃): 1.98 (s, 3H, AcO), 2.03 (s, 3H, AcO), 2.10 (s, 3H, AcO), 2.15 (s, 3H, AcO), 3.39 (t, 2H, $J_{2,1} = 5.1$ Hz, CH₂N₃), 3.62–3.81 (m, 10H, $5 \times$ CH₂O), 4.03–4.12 (m, 2H, H-5, H-6a), 4.29 (dd, 1H, H-6b, $J_{6b,6a} = 12.4$ Hz, $J_{6b,5} = 5.2$ Hz), 4.87 (d, 1H, H-1, $J_{1,2} = 1.7$ Hz), 5.26 (dd, 1H, H-2, *J*_{2,3} = 3.2 Hz, *J*_{2,1} = 1.7 Hz), 5.28 (t, 1H, H-4, *J*_{4,3} = *J*_{4,5} = 9.9 Hz), 5.35 (dd, 1H, H-3, $J_{3,4}$ = 9.9 Hz, $J_{3,2}$ = 3.3 Hz). ¹³C NMR (CDCl₃): 20.61 (CH₃), 20.63 (CH₃), 20.7 (CH₃), 20.8 (CH₃), 50.7 (CH₂), 60.4 (CH₂), 66.2 (CH), 67.4 (CH₂), 68.4 (CH), 69.1 (CH), 69.6 (CH), 70.0 (2 × CH₂), 70.7 (CH₂), 70.8 (CH₂), 97.7 (CH), 169.7 (C), 169.8 (C), 169.9 (C), 170.6 (C); HRMS (ESI) for C₂₀H₃₁O₁₂N₃Na [M + Na]⁺, calcd 528.1805, found 528.1805 (Figures S1 and S2).

Compound 4

To a solution of **3** (225 mg, 0.445 mmol) in MeOH (4.5 mL), PTSA (77 mg, 0.449 mmol) was added, followed by 10% Pd-C (10% w). The resulting suspension was stirred under H_2 atmosphere (1 atm) for 12 h. The Pd/C was removed by filtration through Celite and



Compound 2

To a solution of peracetylated mannose 1 (2 g, 5.128 mmol, 1 eq) in dry DCM (51 mL) containing oven-dried powdered 4 Å molecular sieves, 2-[2-(2-chloroethoxy)ethoxy]ethanol (2.99 g, 7.692 mmol, 1.5 eq) was added and stirred at 0°C under N_2 atmosphere for the filtrate was evaporated under reduced pressure to produce the ammonium salt, as confirmed by ¹H NMR.

The crude of the reaction was dissolved in a 1:1 mixture of $H_2O/MeOH$ (20 mL), to which Amberlite IRN78 basic resin was



added. After 3 h of stirring at 20°C, the reaction mixture was filtrated and evaporated under reduced pressure to yield deprotected amine 4, a colorless oil (137 mg, 0.440 mmol, 99%), which was used in the next step without further purification. ¹H NMR (MeOD): 2.81 (m, 2H, CH₂NH₂), 2.15–3.9 (m, 16H, 5 × CH₂O, H-2, H-3, H-4, H-5, H-6a, H-6b), 4.81 (d, 1H, H-1, $J_{1,2} = 1.7$ Hz). ¹³C NMR (MeOD): 41.9 (CH₂), 62.9 (CH₂), 67.7 (CH), 68.6 (CH₂), 71.3 (CH₂), 71.4 (CH₂), 71.6 (CH), 72.1 (CH), 72.5 (CH), 72.8 (CH₂), 74.6 (2 × CH₂), 101.7 (CH); HRMS (ESI) for C₁₂H₂₆NO₈ [M + H]⁺, calcd 312.1658, found 312.1651 (Figures S3 and S4).

Compound 5

To a solution of 4 (138 mg, 0.444 mmol) in dry DMF (4.0 mL), p-phenylene diisothiocyanate (426 mg, 0.219 mmol, dissolved in 4.0 mL of dry DMF) was added dropwise under N2 atmosphere using a syringe pump over the course of 1 h. After 2 h of stirring at 20°C, the solvent was evaporated under reduced pressure and the residue was purified by flash chromatography (SiO₂, DCM/MeOH: 100/0 \rightarrow 80/20) to yield the thioisocyanate 5 (98 mg, 0.195 mmol, 44%), a light-yellow solid. ¹H NMR (DMSO-d6): 3.29–3.72 (m, 18H, 5 \times CH₂O, CH2NH, H-2, H-3, H-4, H-5, H-6a, H-6b), 4.45 (t, 1H, OH, $J_{\rm OH,H}$ = 5.9 Hz), 4.58 (d, 1H, OH, $J_{\rm OH,H}$ = 5.9 Hz), 4.64 (d, 1H, H-1, $J_{1,2} = 1.4$ Hz), 4.73 (d, 1H, OH, $J_{OH,H} = 4.5$ Hz), 4.76 (d, 1H, OH, $J_{\rm OH,H}$ = 4.7 Hz), 7.37 (d, 2H, $J_{\rm H,H}$ = 8.9 Hz), 7.59 (d, 2H, $J_{\rm H,H}$ = 8.9 Hz), 7.93 (bs, 1H, NH), 9.79 (bs, 1H, NH). ¹³C NMR (DMSO-d6): 43.6 (CH₂), 61.3 (CH₂), 65.7 (CH₂), 67.0 (CH), 68.5 (CH₂), 69.5 (CH₂), 69.6 (CH₂), 69.8 (CH₂), 70.3 (CH), 70.9 (CH), 73.9 (CH), 100.0 (CH), 123.1 (2 × CH), 124.7 (C), 126.2 (2 × CH), 132.6 (C), 139.3 (C), 180.3 (C). HPLC: t_R = 8.17 min, mobile phase consisting of water (solvent A) and acetonitrile (solvent B). Gradient mix: 0.0 min [95% A; 5% B], 20.0 min [77% A; 23% B]. Flowrate: 1 mL/min; HRMS (ESI) for C₂₀H₂₉N₃O₈S₂Na [M + Na]⁺, calcd 526.1294, found 526.1292 (Figures S5-S7).

Figure 5. RQ/vg ratio in neuroretinas and RPE of rats Individual ratio of GFP mRNA levels (relative quantity [RQ]) to vector copy number expressed as viral genome/diploid genome number (vg/dg), measured 4 weeks post-injection in the left and right neuroretinas (A) and in the RPE (B) of rats injected with vehicle, AAV2, or AAV2 + 5 (3^{E6}) (second coupling). Results are expressed as mean \pm SEM. Mann-Whitney test was used for statistical analyses: **p < 0.01.

Compound 6

To a solution of the acetyl glycoside, 1 (500 mg, 1.282 mmol) and 2-(2-ethoxyethoxy)ethanol (258 mg, 1.923 mmol) in dry DCM (13 mL), 4 Å MS, was added. The resulting solution was stirred at 20°C under N₂ atmosphere. After 30 min, F₃B·OEt₂ (791 µL, 6.410 mmol) was added dropwise at 0°C under N₂ atmosphere. The reaction mixture was stirred for 30 min at 0°C and warmed to 20°C. After 12 h of stirring, the MS was filtered and the solvent evaporated. The residue was redissolved in DCM, washed respectively with saturated aqueous NaHCO3, water, and brine, dried over MgSO₄, filtered, and evaporated. The residue was purified by flash chromatography (SiO₂, Cy/AcOEt 70/30) to yield the derivative 7 (430 mg, 0.926 mmol, 72%), a colorless oil. ¹H NMR (CDCl₃): 1.20 (t, 3H, CH₃CH₂O, J = 7.0 Hz), 1.97 (s, 3H, AcO), 2.02 (s, 3H, AcO), 2.09 (s, 3H, AcO), 2.14 (s, 3H, AcO), 3.52 (q, 2H, CH₃CH₂O, J = 7.0 Hz), 3.56–3.86 (m, 8H, 4 × CH₂O), 4.08 (m, 2H, H-5, H-6a), 4.28 (dd, 1H, H-6b, J_{6b,6a} = 12.6 Hz, $J_{6b,5} = 5.3$ Hz), 4.89 (d, 1H, H-1, $J_{1,2} = 1.5$ Hz), 5.25 (dd, 1H, H-2, $J_{2,3} = 3.3$ Hz, $J_{2,1} = 1.8$ Hz), 5.27 (t, 1H, H-4, $J_{4,3} = J_{4,5} =$ 9.7 Hz), 5.35 (dd, 1H, H-3, $J_{3,4}$ = 10.0 Hz, $J_{3,2}$ = 3.4 Hz). ¹³C NMR (CDCl₃): 15.1 (CH₃), 20.6 (CH₃), 20.7 (CH₃), 20.8 (CH₃), 62.4 (CH₂), 66.1 (CH), 66.6 (CH₂), 67.4 (CH), 68.4 (CH), 69.1 (CH), 69.6 (CH₂), 69.8 (CH₂), 70.0 (CH₂), 70.8 (CH₂), 97.7 (CH), 169.7 (C), 169.8 (C), 170.0 (C), 170.1 (C); HRMS (ESI) for C₂₀H₃₂O₁₂Na [M + Na]⁺, calcd 487.1791, found 487.1784 (Figures S8 and S9).

Compound 7

The acetylated sugar **6** (310 mg, 0.667 mmol) was dissolved in a 1:1 mixture of $H_2O/MeOH$ (12 mL), to which Amberlite IRN78 basic resin was added. After 3 h of stirring at 20°C, the reaction mixture was filtrated and evaporated under reduced pressure to yield the final 7 (195 mg, 0.659 mmol, 99%), a white solid. ¹H NMR (MeOD): 1.19





Figure 6. Ocular fundus in NHPs

Kinetics of GFP expression after subretinal delivery of vehicle, AAV2, or AAV2 + **5** ($3^{E}6$), using *in vivo* fundus imaging. The green color becomes quickly saturated for AAV2 + **5** ($3^{E}6$) as the expression of GFP is very high and the sensitivity of image acquisition on the retinophotograh was fixed for all retinal photographs from 1 wpi. Note that retinal structure was preserved as normal on retinophotographs with lower sensitivity and as seen by OCT (data not shown). wpi, weeks post-injection.

cells were harvested 48 h after transfection and treated with Triton-1% and benzonase (25 U/mL) for 1 h at 37°C. The resulting bulk was subjected to freeze-thaw cycles to release vector particles. The cellular debris was removed by centrifugation at 2,500 rpm for 15 min. Cell lysates were precipitated with PEG overnight and clarified by centrifugation at 4,000 rpm for 1 h. The precipitates were then incubated with benzonase for 30 min at 37°C and collected after centrifugation at 10,000 \times g for 10 min at 4°C. Vectors were purified by double cesium chloride (CsCl) gradient ultracentrifugation. The viral suspension was then subjected to four successive rounds of dialysis with mild stirring in a Slide-a-Lyzer cassette (Pierce) against dPBS (containing Ca²⁺ and Mg^{2+}).

Coupling and purification

AAV2-GFP ($1^{E}12$ vg, 2.49 nmol) was added to a solution of TBS buffer (pH 9.3) containing 5 or 7 at different molar ratios ($3^{E}5$ or $3^{E}6$ eq), as stated in the respective results sections, and incubated for 4 h at RT. The solutions containing the vectors were then dialyzed against

(t, 3H, **CH**₃CH₂O, *J* = 7.0 Hz), 3.54 (q, 2H, CH₃**CH**₂O, *J* = 7.0 Hz), 3.56–3.86 (m, 14H, $4 \times$ **CH**₂O, H-2, H-3, H-4, H-5, H-6a, H-6b), 4.79 (d, 1H, H-1, *J*_{1,2} = 1.7 Hz). ¹³C NMR (MeOD): 15.4 (CH₃), 62.9 (CH₂), 67.6 (CH), 67.8 (CH₂), 68.6 (CH), 70.9 (CH), 71.4 (CH), 71.6 (CH₂), 72.1 (CH₂), 72.5 (CH₂), 74.6 (CH₂), 101.7 (CH). HPLC: t_R = 2.19 min, mobile phase consisting of water (solvent A) and acetonitrile (solvent B). Gradient mix: 0.0 min [95% A; 5% B], 20.0 min [77% A; 23% B]. Flowrate: 1 mL/min; HRMS (ESI) for C₁₂H₂₄O₈Na [M + Na]⁺, calcd 319.1369, found 319.1371 (Figures S10–S12).

AAV2 production and purification

AAV2 vectors were produced from two plasmids: (1) pHelper, PDP2-KANA encoding AAV Rep2-Cap2 and adenovirus helper genes (E2A, VA RNA, and E4) and (2) the pVector ss-CAG-eGFP containing the ITRs. All vectors were produced by transient transfection of HEK293 cells using the calcium phosphate-HeBS method. AAV2-transfected dPBS + 0.001% Pluronic to remove free molecules that had not bound to the AAV capsid.

Titration of AAV vector genomes

A total of 3 μ L of AAV was treated with 20 units of DNase I (Roche, no. 04716728001) at 37°C for 45 min to remove residual DNA in vector samples. After treatment with DNase I, 20 μ L of proteinase K (20 mg/mL; Macherey-Nagel, no. 740506) was added and the mixture incubated at 70°C for 20 min. An extraction column (NucleoSpinRNA Virus) was then used to extract DNA from purified AAV vectors.

Real-time quantitative PCR (qPCR) was performed with a StepOnePlus Real-Time PCR System Upgrade (Life Technologies). All PCRs were performed with a final volume of 20 μ L, including primers and probes targeting the ITR2 sequence, PCR Master Mix (TaKaRa), and 5 μ L of template DNA (plasmid standard or sample



Figure 7. Fluorescence intensity of GFP expression



DNA). qPCR was carried out with an initial denaturation step at 95° C for 20 s, followed by 45 cycles of denaturation at 95° C for 1 s and annealing/extension at 56° C for 20 s. Plasmid standards were generated with seven serial dilutions (containing 10^{8} to 10^{2} plasmid copies), as described by D'Costa et al.⁴²

Western blot and silver staining

All vectors were denatured at 100°C for 5 min using Laemmli sample buffer and separated by SDS-PAGE on 10% Tris-glycine polyacrylamide gels (Life Technologies). Precision Plus Protein All Blue Standards (Bio-Rad) were used as a molecular weight size marker. After electrophoresis, gels were either silver stained (PlusOne Silver Staining Kit, Protein; GE Healthcare) or transferred onto nitrocellulose membranes for western blot. After transferring the proteins to nitrocellulose membrane using a transfer buffer (25 mM Tris/192 mM glycine/0.1 [w/v] SDS/20% MeOH) for 1 h at 150 mA in a Trans-Blot SD Semi-Dry Transfer Cell (Bio-Rad), the membrane was saturated for 2 h at RT with 5% semi-skimmed milk in PBS-Tween (0.1%) or with 1% gelatin, 0.1% Igepal in PBS-Tween (0.01%). After saturation, the membrane was probed with the corresponding antibody (anti-capsid polyclonal, B1 monoclonal, or anti-fluorescein-AP) and FITC-Concanavalin A lectin (mannose detection) overnight at 4°C. Three washes (15 min at RT) with PBS-Tween (0.1%) were performed between each stage to remove unbound reagents. Bands were visualized by chemiluminescence using AP- or horseradish peroxidase-conjugated secondary antibodies and captured on X-ray film.

DLS

DLS was performed using a Malvern Zetasizer Nano ZS. Calibration was controlled beforehand using a 30 and 300 nm solution of Nanosphere Size Standard. A volume of 50 μ L of each vector was placed in a specific cuvette (DTS0118; Malvern) and analyzed by volume.

AAV infectious titer measurement

The infectivity of each sample was measured as follows. HEK293 or HeLa cells were seeded in 2 mL DMEM growth medium in 6-well cul-

ture plates at a density of 10^6 cells/well. Cells were then incubated overnight at 37° C to reach 50% confluence. The viral stock was then diluted 10-fold by serial dilution. Next, 2 µL of each dilution was added to separate wells in the six-well plates. Plates were then incubated at 37° C for 24 h. The infectivity of the AAV2-GFP control was measured immediately upon thawing of the sample. The same procedure was used for mannose particles. AAV-GFP-infected cells were detected by fluorescence microscopy.

The transducing unit (TU) titer was calculated using the following formula:

 $TU / mL = (4,040 \times NGFP \times dilutions \times 1,000) / V$

where NGFP is the mean number of GFP-positive cells per well and V is the volume (in μ L) of vector used to infect cells.

Animal care and welfare

Experiments were performed on 33 rats (Sprague-Dawley) and 3 male cynomolgus monkeys (*Macaca fascicularis*) provided by BioPrim (Baziege, France). Animals were euthanized 1 month after AAV injection. Research was conducted at the Boisbonne Centre (ONIRIS, Nantes-Atlantic College of Veterinary Medicine, Nantes, France) under authorization no. H44273 from the Departmental Direction of Veterinary Services (Loire-Atlantique, France). All animals were handled in accordance with the Guide for the Care and Use of Laboratory Animals. Experiments involving animals were conducted in accordance with the guidelines of the Animal Experimentation Ethics Committee of Pays de Loire (France) and the Ministry of Higher Education and Research. Animals were sacrificed by intravenous injection of pentobarbital sodium (Dolethals; Vetoquinol) in accordance with approved protocols.

Subretinal injections

Injection in rats

Rats were anesthetized by inhalation of isoflurane gas followed by intramuscular injection of ketamine/xylazine (Imalgène1000/Rompun 2%). Pupils were dilated with 0.5% tropicamide and 10% neosynephrine eye drops. Local anesthesia was achieved using 0.4% oxybuprocaine eye drops. The ocular surface was disinfected with 5% vetedine and rinsed 3 times with Ocryl. Under an operating microscope a trans-scleral/transchoroidal tunnel incision was made using surgical nylon suture (Ethilon 10/0) and the AAVs delivered to the subretinal space by injection using a 33G needle and a 5-mL Hamilton syringe. Slow injection was performed at a rate of 2 μ L/min while monitoring using a binocular microscope. A volume of 2.5 μ L was injected with 1/1,000 of fluorescein dye to monitor bleb size. Injection success was confirmed by fundus autofluorescence imaging immediately after injection by SD-OCT (Spectralis HRA+OCT imaging system; Heidelberg Engineering).

Injection in NHPs

NHPs were anesthetized by isoflurane gas inhalation. Pupils were dilated with atropine 0.3%, tropicamide 0.5%, and neosynephrine



10% eye drops. Local anesthesia was achieved with oxybuprocaine 0.4% eye drops. The ocular surface was disinfected with vetedine 5% and rinsed three times with Ocryl. After lateral canthotomy, a 25-gauge 3-port vitrectomy was performed with posterior vitreous detachment (Vitreotom Accurus; Alcon France). Retinal bleb formation was induced by subretinal injection of viral vector (150 µL) along the temporal superior vascular arcade with a 41-gauge subretinal injection cannula. At the end of the procedure, fluid-air exchange of the vitreous cavity was performed. Sclerectomies and the lateral canthotomy were sutured with Vicryl 7/0 (absorbable). Subconjunctival injection of 0.5 mL methylprednisolone (40 mg/mL) was performed.

NHP 3 $(AAV2 + 5 (3^{E}6))$



Figure 8. Retinal tropism of AAV2 + 5 (3^E6) and AAV2 vectors in flat-mount retinas from NHPs

Images are taken from confocal slices of each retinal layer and show GFP expression (green), nuclear labeling (purple), and immunostaining with LM opsin to identify outer segments (red). Original magnification, ×60. Sensitivity of image acquisition was set on the retina with injection of AAV2, at the minimum to allow the visualization of the whole transduced retina. Images shown on this figure correspond to the temporal retina of both NHP2 and NHP3, where the GFP expression was maximal.

Retinal imaging

For evaluation of retinal structure, SD-OCT was performed before AAV injection and after euthanasia using the Spectralis HRA+OCT imaging system (Heidelberg Engineering). Eye fundoscopy was performed before injection and 1, 2, 3, 4, and 6 weeks post-injection to monitor GFP expression over time, using a Canon UVI retinal camera connected to a digital imaging system (LHEDIOPH 1600; AMS Ophta). To quantify the GFP level of every fundus, a diagonal line (left bottom to top right) was plotted on images exported from the Canon UVI retinal camera. Intensity level in GFP (green channel) was measured averaged over a width of line of 21 pixels (average of 250 points on the diagonal line).

Whole-mount immunofluorescence in NHP retinas

Eyes of NHPs were enucleated and fixed for 1 h in 4% paraformaldehyde in phosphate-buffered saline (PBS) solution at RT. Next, the anterior part and the lens was removed and the neuroretina separated from the RPE-choroid-sclera using fine forceps. Retinas were then washed 3 times in PBS at RT and incubated with polyclonal rabbit LM opsin and S opsin antibody (1/300; Vector Laboratories, Peterborough, UK)

for 48 h at 4°C. After 3 washes in PBS, retinas were incubated for 2 h at RT with 546-conjugated anti-rabbit IgG antibody (1/300; Life Technologies, Saint Aubin, France). Cell nuclei were labeled with Draq5 (1/500; Biostatus, Leicestershire, UK) for 2 h at RT. Retinas were subsequently washed in PBS, mounted on slides with coverslips in Prolong Gold anti-fade reagent (Life Technologies), and stored at RT before microscopic analysis. Flat-mounted neuroretinas were examined using a laser scanning confocal microscope (Nikon A1RSi). Three-dimensional digital images were collected at 20× (NA 0.75) and $60 \times$ (NA 1.4) using NIS-Elements confocal software and appropriate fluorescence filters for laser excitation at 488, 647, and 546 nm.

Immunohistochemistry GFP staining in rats

Rat eyes were enucleated, fixed for 5 h in Bouin's solution, dehydrated in successive alcohol solutions, and embedded in paraffin. Entire retinas were sliced into 5-µm sections. Paraffin tissue sections were deparaffinized and rehydrated, and antigen was unmasked by boiling the section in ready-to-use citrate-based solution (pH 6.0). Sections were permeabilized for 10 min at RT in PBS containing 0.2% Triton X-100. Non-specific activity was blocked by incubating the sections for 45 min at RT in 2% goat serum in PBS and 5% BSA in PBS. Sections were incubated overnight at 4°C in primary antibody diluted in PBS (1:100 monoclonal mouse anti-GFP; 632380, Clontech). After washing with PBS, sections were incubated for 1 h at RT with goat anti-mouse Alexa 555-conjugated antibody (1:300; A21434, Life Technologies). After counterstaining with hematoxylin, slides were mounted under coverslips in Prolong Gold anti-fade reagent (Life Technologies). Stained slides were scanned using the Hamamatsu NanoZoomer (Bacus Laboratories, Chicago, IL) and analyzed using NDP.View2 software. Mayer's hemalum staining images and the corresponding fluorescence image were merged using Fiji software (http://fiji.sc).

Absolute quantification of vector genomes by qPCR in rat tissue samples

Samples (neuroretinas + RPE/choroid) were obtained just after sacrifice in conditions that minimized cross contamination and avoided qPCR inhibition, as described by Le Guiner et al.⁴³ Samples were snap-frozen in liquid nitrogen and stored at \leq -70°C before DNA extraction. Genomic DNA (gDNA) was extracted from the whole retinal pigment epithelium using the Nucleospin Tissue Kit (Macherey Nagel) and TissueLyserII (QIAGEN) according to the manufacturer's instructions. qPCR was performed on a StepOne Plus Real-Time PCR System (Applied Biosystems, Thermo Fisher Scientific) using 50 ng of gDNA in duplicate. Vector genome copy number was determined using the following primer/probe combination, designed to amplify a specific region of the GFP sequence present in the transgene (forward, 5'-ACTACAACAGC CACAACGTCTATATCA-3'; reverse, 5'-GGCGGATCTTGAAGTT CACC-3'; probe, 5'-FAM- CCGACAAGCAGAAGAACGGCAT CA-TAMRA-3'). Endogenous gDNA copy number was determined using a primer/probe combination designed to amplify the rat Hprt1 gene (forward, 5'-GCGAAAGTGGAAAAGCCAAGT-3'; reverse, 5'- GCCACATCAACAGGACTCTTGTAG -3'; probe, 5'-FAM- CA AAGCCTAAAAGACAGCGGCAAGTTGAAT -TAMRA-3'). For each sample, cycle threshold (Ct) values were compared with those obtained with different dilutions of linearized standard plasmids (containing either the eGFP expression cassette or the rat Hprt1 gene). The absence of qPCR inhibition in the presence of gDNA was determined by analyzing 50 ng gDNA extracted from tissue samples from a control animal and spiked with different dilutions of standard plasmid. Results are expressed as vg/dg. The LLOQ of our test was 0.002 vg/dg. GraphPad Prism 9 software was used for statistical analysis. The non-parametric Mann-Whitney test was used.

Relative quantification of GFP expression by real-time qPCR in rat tissue samples

RNA was extracted from neuroretinas and whole RPE, using the Nucleospin RNA kit (Macherey Nagel) and TissueLyserII (QIAGEN) according to the manufacturer's instructions. One thousand nanograms of total RNA was treated with RNAse-free DNAse I (ezDNAse from Thermo Fisher Scientific) and then reverse transcribed using SuperScript IV Vilo reverse transcriptase (Thermo Fisher Scientific) in a final volume of 20 µL. qPCR analysis was then performed on cDNA (diluted 1/20) using the same GFP primers and probe as for the quantification of transgene copy number by qPCR. Hprt1 messenger was also amplified as an endogenous control. For each RNA sample, the absence of DNA contamination was confirmed by analysis of "cDNA-like samples" obtained without adding reverse transcriptase to the reaction mix. The absence of qPCR inhibition in the presence of cDNA was determined by analyzing cDNA obtained from RPE from a control animal, spiked with different dilutions of a standard plasmid. For each RNA sample, Ct values were compared with those obtained with different dilutions of standard plasmids (containing either the GFP expression cassette or the *Hprt1* gene). Results are expressed in RQ: $RQ = 2-\Delta Ct = 2-(Ct)$ target - Ct endogenous control). The LLOQ of our test was RQ = 9^E-4. GraphPad Prism 9 software was used for statistical analysis. Non-parametric Mann-Whitney test was used. Samples were considered significantly different if p < 0.05, p < 0.01, or p < 0.001.

Absolute quantification of vector genomes by qPCR in NHP tissue samples

Samples (liver, spleen, occipital cortex, and chiasma optic) were obtained just after sacrifice under conditions that minimized cross contamination and avoided qPCR inhibition, as described by Le Guiner et al.43 Samples were snap-frozen in liquid nitrogen and stored at \leq -70°C before DNA extraction. gDNA was extracted using the Gentra puregene kit and TissueLyserII (both from QIAGEN) according to the manufacturer's instructions. qPCR analyses were conducted on a StepOne Plus Real-Time PCR System (Applied Biosystems, Thermo Fisher Scientific) using 50 ng of gDNA in duplicate. Vector genome copy number was determined using the same GFP-specific primer/ probe combination as used for rat tissue. Endogenous gDNA copy number was determined using a primer/probe combination designed to amplify the macaque Eglobin gene (forward, 5'-TGGCAAGGAGTT CACCCCT-3'; reverse, 5'-AATGGCGACAGCAGACACC-3'; probe, 5'-FAM- TGCAGGCTGCCTGGCAGAAGC -TAMRA-3'). For each sample, Ct values were compared with those obtained with different dilutions of linearized standard plasmids (containing either the eGFP expression cassette or the macaque Eglobin gene). The absence of qPCR inhibition in the presence of gDNA was determined by analyzing 50 ng of gDNA extracted from tissue samples from a control animal and spiked with different dilutions of standard plasmid. Results are expressed as vg/dg. The LLOQ of our test was 0.001 vg/dg.

DATA AND CODE AVAILABILITY

All relevant data are available from the authors on request and are included in the article.

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10. 1016/j.omtm.2024.101187.

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DECLARATION OF INTERESTS

M.M., D.D., and E.A. are inventors on a patent including the technology described in this manuscript. A.G., N.B., and G.M.L. are employees of Coave Therapeutics.

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