Long-term Preservation of Cone Photoreceptors and Restoration of Cone Function by Gene Therapy in the Guanylate Cyclase-1 Knockout (GC1KO) Mouse

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PURPOSE. The authors previously showed that subretinal delivery of AAV5 vectors containing murine guanylate cyclase-1 (GC1) cDNA driven by either photoreceptor-specific (hGRK1) or ubiquitous (smCBA) promoters was capable of restoring cone-mediated function and visual behavior and preserving cone photoreceptors in the GC1 knockout (GC1KO) mouse for 3 months. Here, the authors compared therapy conferred by the aforementioned vectors to that achieved with the highly efficient capsid tyrosine mutant AAV8(Y733F) being the most efficient. Electroretinographic responses were clearly measurable out to 1 year after injection. AAV-mediated expression of GC1 was found exclusively in photoreceptors out to 15 months after injection. Cone function was restored with all vectors tested, with AAV8(Y733F) being the most efficient. Electoretinographic responses were clearly measurable out to 1 year after treatment. AAV-mediated expression of GC1 was found exclusively in photoreceptors out to 15 months after injection. Cones were preserved for at least 11 months after treatment. AAV5- and AAV8(Y733F)-delivered vector genomes were recovered primarily from optic nerve of the treated eye and, in only instance, from brain (1 of 20 samples).

RESULTS. Cone function was restored with all vectors tested, with AAV8(Y733F) being the most efficient. Electoretinographic responses were clearly measurable out to 1 year after treatment. AAV-mediated expression of GC1 was found exclusively in photoreceptors out to 15 months after injection. Cones were preserved for at least 11 months after treatment. AAV5- and AAV8(Y733F)-delivered vector genomes were recovered primarily from optic nerve of the treated eye and, in only instance, from brain (1 of 20 samples).

CONCLUSIONS. The authors demonstrate for the first time that long-term therapy (~1 year) is achievable in a mammalian model of GC1 deficiency. These data provide additional justification for the development of an AAV-based gene therapy vector for the clinical treatment of Leber congenital amaurosis-1. (Invest Ophtalmol Vis Sci. 2011;52:7098–7108) DOI: 10.1167/iovs.11-7867

Retinal guanylate cyclase-1 (GC1) encoded by GUCY2D (Gucy2e in mouse) is expressed in the outer segments of rod and cone photoreceptors of human, monkey, and mouse retinas. 1–5 GC1 plays a vital role in light-dark and recovery cycles, anchoring, through cGMP, the feedback loop linking intracellular calcium levels and the polarization state of photoreceptors. 4–8 Like other membrane guanylate cyclases, it contains an N-terminal signal sequence, an extracellular domain, a single transmembrane domain, a kinase-like homology domain, a dimerization domain, and a C-terminal catalytic domain and is present likely as homodimeric dimers. 9 Mutations in GUCY2D are associated with recessive Leber congenital amaurosis-1 (LCA1) as well as dominant and recessive forms of cone-rod dystrophy, CORD6 and CORD, respectively. 10–16 LCA1 is a severe, early-onset, autosomal recessive blinding disorder characterized by extinguished electroretinogram, which precedes photoreceptor degeneration. 17–18 CORD6 is a dominant disorder characterized by progressive degeneration of photoreceptors beginning with cones causing early loss of visual acuity and color vision followed by degeneration of rods leading to progressive night blindness and peripheral visual field loss. 12,13 CORD6 mutations are restricted to the dimerization domain and generally cause an increase in guanylate cyclase activating protein (GCAP)–mediated activation of GC1. 19–21 A recently found recessive CORD-causing mutation is located in the catalytic domain of GC1 and is thought to reduce overall enzyme function. 16 LCA1-causing mutations are distributed throughout GC1. 22 These mutations alter enzyme structure and stability, may impact retrograde transport of other peripheral membrane-associated proteins, and are frequently null. 22

The GC1 knockout (GC1KO) mouse carries a null mutation in Gucy2e, the murine homolog of GUCY2D. Like LCA1 patients, loss of cone function in this model precedes cone degeneration. 22 Rods retain 30% to 50% of their function and do not degenerate because of the presence of guanylate cyclase-2 (GC2), another functional guanylate cyclase in murine photoreceptors.9,22–25 We have previously shown that subretinal injection of serotype 5 adeno-associated viral (AAV5) vectors containing the murine GC1 cDNA driven by either the photoreceptor-specific human rhodopsin kinase (hGRK1) or the ubiquitous (smCBA) promoter were capable of restoring cone-mediated function and visual behavior and preserving cone photoreceptors in the GC1KO mouse for 3 months. 26 In the present study, we evaluated whether AAV-mediated gene
Electroretinographic Analysis

Subretinal Injections

GC1KO mice between postnatal day (P) 14 and P25. Our findings demonstrate for the first time that long-term therapy (1 year) is achievable in a mammalian model of GC1 deficiency. We also used vector genome biodistribution and tissue distribution of AAV8/Y733F vectors. These findings have direct bearing on the development of an AAV-based gene therapy protocol for both rod and cone degenerative diseases.
vector genome copies were established according to previously described methods, with minor modifications that can be found in the Supplementary Methods.26 Retinas were immediately dissected and frozen in liquid nitrogen. Optic nerves were dissociated from the eyes, fixed in 4% paraformaldehyde overnight at 4°C, immersed in 30% sucrose for 2 hours at 4°C, and then quick frozen in cryostat compound (Tissue Tek OCT 4588; Sakura Finetek USA, Inc., Torrance, CA) in a bath of dry ice/ethanol. Optic nerves were sectioned at 10 μm and stained according to previously described methods.28 Retinas were homogenized in 350 μL lysis buffer (Buffer RLT; RNeasy Protect Mini Kit; Qiagen Inc., Valencia, CA) plus BME for 45 seconds. Samples were centrifuged, and the lysate was split in half (one half designated for genome recovery and the other half for RNA extraction).34 Genome recovery was performed as described. RNA extraction was performed with a stabilization and purification kit (RNeasy Protect Mini Kit; Qiagen Inc.). RNA was reverse transcribed (Script cDNA synthesis kit; Bio-Rad Laboratories) and used in real-time PCR (iQ SYBR Green Supermix and Mq real-time PCR detection system interfaced with iCycler thermal cycler; Bio-Rad Laboratories) to measure the following retinal-specific mRNAs: GC1, GCAP1, cone transducin alpha (GNAT2), rod CGMP-specific 3′,5′ cyclic phosphodiesterase subunit alpha (PDE6a), and the housekeeping gene glyceraldehyde 3-phosphate dehydrogenase (GAPDH). Primer sequences and a detailed description of target validation are provided in the Supplementary Methods.26

RESULTS

mRNA Quantification by Real time-PCR, Retinal Genome Recovery, and Optic Nerve Immunohistochemistry

Treated eyes with optic nerves attached were harvested from GC1KO mice 1 year after treatment with either AAV8(Y733F)-hGRK1-mGC1 or AAV5-smCBAm-mGC1 and an age-matched, untreated GC1+/+ mouse. Retinas were immediately dissected from the eye and snap frozen in liquid nitrogen. Optic nerves were dissociated from the eyes, fixed in 4% paraformaldehyde overnight at 4°C, immersed in 30% sucrose for 2 hours at 4°C, and then quick frozen in cryostat compound (Tissue Tek OCT 4588; Sakura Finetek USA, Inc., Torrance, CA) in a bath of dry ice/ethanol. Optic nerves were sectioned at 10 μm and stained according to previously described methods.28 Retinas were homogenized in 350 μL lysis buffer (Buffer RLT; RNeasy Protect Mini Kit; Qiagen Inc., Valencia, CA) plus BME for 45 seconds. Samples were centrifuged, and the lysate was split in half (one half designated for genome recovery and the other half for RNA extraction).34 Genome recovery was performed as described. RNA extraction was performed with a stabilization and purification kit (RNeasy Protect Mini Kit; Qiagen Inc.). RNA was reverse transcribed (Script cDNA synthesis kit; Bio-Rad Laboratories) and used in real-time PCR (iQ SYBR Green Supermix and Mq real-time PCR detection system interfaced with iCycler thermal cycler; Bio-Rad Laboratories) to measure the following retinal-specific mRNAs: GC1, GCAP1, cone transducin alpha (GNAT2), rod CGMP-specific 3′,5′ cyclic phosphodiesterase subunit alpha (PDE6a), and the housekeeping gene glyceraldehyde 3-phosphate dehydrogenase (GAPDH). Primer sequences and a detailed description of target validation are provided in the Supplementary Methods.26

RESULTS

Long-term, Photoreceptor-Specific GC1 Expression Is Achievable in GC1KO Mice Treated with AAV5 or AAV8(Y733F) Vectors

Immunostaining with an antibody directed against GC1 (green) revealed that AAV-vectorized therapeutic protein expression persisted exclusively in photoreceptors of treated GC1KO mice for a significant fraction of the animal’s lifetime; AAV8(Y733F)-hGRK1-mGC1 for at least 7 months, AAV5-smCBAm-mGC1 for at least 10 months, and AAV5-hGRK1-mGC1 for at least 15 months (Figs. 1A, 1B). GC1 expression was limited to the outer segments of rods and cones treated with AAV8(Y733F)-hGRK1-mGC1 vector, whereas it was found in both outer segments and more rarely in photoreceptor cell bodies of eyes treated with AAV5-smCBAm-mGC1. GC1 expression was absent from the untreated GC1KO retina (Figs. 1A, 1B). Orientation of eyes before kill allowed for the evaluation of GC1 expression in similar areas of treated or untreated retinas (inferior retina), with care taken to avoid analysis in the far periphery. Two examples of retinal thinning were observed. The first was a GC1KO retina treated with AAV5-smCBAm-mGC1 (4.69 ± 10^0 total vector genomes delivered) (Fig. 1A). The outer nuclear layer (ONL) was slightly thinned relative to that seen in naive GC1KO or GC1+/+ control retinas (both 8 months of age). The second was a GC1KO
Cones remained in the superior retina. Wholemount analysis of a cross-section taken from the inferior retina of an untreated AAV8(Y733F)-hGRK1-mGC1 (7 months after injection) vector revealed increased GC1-positive outer segments found in the former example was due to overall reduction in photoreceptor outer segments after injection-related damage.

**Long-term Cone Photoreceptor Survival Is Achieved by AAV-Vectored GC1**

Cone photoreceptors in treated and untreated GC1KO mice as well as GC1+/+ controls were identified by staining for mouse cone arrestin (red). Here we show that cone photoreceptor densities were markedly reduced in untreated GC1KO retinas by 8 months of age (Fig. 1) and confirm previous reports that cones are lost in a topographically specific manner in this animal. We previously showed that average photopic b-wave amplitudes in AAV-mGC1–treated mice were partially restored at 4 weeks after injection and remained stable for 3 months.26 We also compare, out to 6 months, cone responses in mice treated with AAV5-hGRK1-mGC1 to these cohorts. Cone responses were stable over time and were significantly revealed the 12-month-old untreated retina exhibited sparse cone density, with residual cones found exclusively in superior retinal regions whereas the partner P14-treated retina retained much higher cone density throughout, with the exception of a small patch of temporal retina that likely was not exposed to vector during subretinal injection and therefore did not contain transgene product (Fig. 2). Quantification of cone photoreceptors revealed significant differences in cone cell densities in the inferior (P < 0.001), central (P < 0.001), and superior (P < 0.001) retinal regions of treated and untreated GC1KO retinas. Average cone cell densities per square millimeter in the treated GC1KO retina were 1597 (inferior), 1432 (central), and 1790 (superior). Average cone cell densities per square millimeter in the untreated GC1KO retina were 2 (inferior), 7 (central), and 637 (superior). These results show that AAV-mGC1 treatment is capable of preserving cone photoreceptors for at least 11 months after treatment and suggest this that preservation of cones is possible over the lifetime of the animal.

**Long-term Restoration of Photoreceptor Function (ERG) Is Achieved in AAV-Treated GC1KO Mice**

We previously showed that average photopic b-wave amplitudes in AAV-mGC1–treated mice were partially restored at 4 weeks after injection and remained stable for 3 months.26 Here we statistically compare cone-mediated responses out to 9 months after treatment in cohorts of mice treated with AAV5 vector containing hGRK1 or smCBA promoters, the identical vectors used in our previous 3-month study.26 We also compare, out to 6 months, cone responses in mice treated with AAV8(Y733F)-hGRK1-mGC1 to these cohorts. Cone responses were stable over time and were significantly lower than the respective untreated AAV8(Y733F)-hGRK1-mGC1 and AAV5-smCBA-mGC1 treated retina (Fig. 2).
higher than responses generated from untreated contralateral controls ($P < 0.001$), suggesting that restoration of cone function is possible over the lifetime of the animal (Fig. 3). Consistent with our previous report, the level of restoration achieved after delivery of the photoreceptor-specific promoter (hGRK1)-containing AAV5 vector was not significantly different from that achieved with the ubiquitous promoter (smCBA)-containing AAV5 vector at any post-treatment time point. Beyond the period of statistical significance, a higher average number of vector genomes in the optic nerves of eyes injected with AAV5-GRK1-mGC1 compared with AAV5-smCBA-mGC1. Only AAV5-hGRK1-mGC1–delivered vector genomes were detected in the right optic nerves of AAV5-treated mice at both 7 and 10 months after injection. At 7 months after injection, vector genomes were also detected in the left optic nerves. At 10 months after injection, AAV5-delivered vector genomes were still detected in right (injected) optic nerve but were absent from both brain hemispheres. AAV8(Y733F)-delivered vector genomes were detected in the left brain at both 4 and 7 months after injection.

Calculating the average ratios of rod b-wave amplitudes in AAV8(Y733F)-hGRK1-mGC1–treated versus untreated eyes resulted in values $>1.0$, with one exception at 4 months after treatment (Fig. 4). Rod b-wave ratios of AAV5-treated versus untreated eyes were only occasionally $>1.0$. Similarly, rod a-wave ratios were consistently $>1.0$ in AAV8(Y733F)-hGRK1-mGC1–treated mice whereas they were, for the most part, $<1.0$ after treatment with either AAV5 vector (Fig. 4). Representative rod-mediated scotopic ERG traces elicited by a $1 \text{ cd} \cdot \text{s/m}^2$ stimulus show AAV8(Y733F)-mediated improvements in rod ERG amplitudes and indicate that aside from the reduced amplitude, treated-eye response kinetics resemble that seen in the GC1+/+ control (Fig. 4).

**Vector Biodistribution**

Optic nerve from injected and un.injected eyes and portions of left and right brain that contained visual pathways were analyzed for the presence of vector genomes. AAV5 vectors were injected in the right eyes of GC1KO mice. Accordingly, vector genomes were detected in the right optic nerve of AAV5-treated mice at both 7 and 10 months after injection. At 7 months after injection, vector genomes were also detected in the left optic nerves of mice injected with AAV5-hGRK1-mGC1. At 10 months after injection, AAV5-delivered vector genomes were still detected in right (injected) optic nerve but were absent from both brain hemispheres. AAV8(Y733F) vector was injected into the left eyes of GC1KO mice. Accordingly, AAV8(Y733F)-delivered vector genomes were detected in the left optic nerves at both 4 and 7 months after injection. At no time point were vector genomes in the AAV8(Y733F)-treated mouse detected in either brain hemisphere. We detected a higher average number of vector genomes in the optic nerves of eyes injected with AAV5-GRK1-mGC1 compared with AAV5-smCBA-mGC1. Only AAV5-hGRK1-mGC1–delivered genomes were detected in brain tissue over the course of our study (1 of 16 samples). As expected, no AAV vector genomes were recovered from any tissue of naive GC1KO control mice (Table 2).

**AAV-mGC1 Treatment Restores Wild-Type Levels of GC1 and GCP1 to Treated GC1KO Retina**

GC1 protein was absent from the untreated eye of the GC1KO mouse. In contrast, at 7 months after injection with AAV8(Y733F)-hGRK1-mGC1, the level of GC1 in the treated eye was comparable to that seen in the GC1+/+ control (Fig. 5A).
Consistent with previous reports that GCAP1 is post-translationally downregulated in the GC1KO mouse, we show that GCAP1 was downregulated in the untreated GC1KO retina relative to the GC1+/+ control (Fig. 5A). However, AAV-mediated delivery of GC1 leads to an upregulation in GCAP1 expression in the treated GC1KO mouse retina. Levels of GCAP1 expression were also comparable to those seen in GC1+/+ controls (Fig. 5A).

**FIGURE 3.** Cone-mediated electroretinograms are restored in AAV-mGC1-treated GC1KO mice over the long term. (A) Representative cone-mediated traces elicited by a 12 cd·s/m² light stimulus from GC1KO eyes treated with AAV5-hGRK1-mGC1 (red line), AAV5-smCBA-mGC1 (green line), or AAV8(Y733F)-hGRK1-mGC1 (black line) or age-matched, untreated GC1+/+ control eyes reveal long-term AAV-mGC1-mediated restoration of cone function. Representative traces generated between 4 months and 1 year after treatment are shown. (B) Maximum cone b-wave amplitudes (those generated at 12 cd·s/m²) were calculated from each mouse and averaged monthly in each treatment group and in age-matched, untreated GC1KO and GC1+/+ controls. Comparisons were made between groups of animals with an n ≥ 3. All AAV treatment groups were statistically compared for 6 months after treatment. AAV5 vector-treated eyes were statistically compared for 9 months after treatment. Error bars represent ±1 SD.
In Treated GC1KO Mice, GC1 mRNA Is Present and GNAT2 mRNA Levels Are Increased Compared with Untreated GC1KO Mice

At 1 year after treatment, levels of GC1 mRNA in treated retinas were approximately 7-fold (AAV5-treated) and 14-fold (AAV8[Y733F]-treated) higher than those seen in the age-matched GC1+/H11001/ control mouse (Fig. 5B). We found that high levels of GC1 mRNA in treated retinas corresponded to recovery of many vector genomes: 1.57 x 10^7 vg/µg DNA for AAV8(Y733F) and 4.7 x 10^6 vg/µg for AAV5. Despite the high levels of GC1 mRNA in treated retinas, no GC1 expression was detected in optic nerves of treated eyes (data not shown). This result further supports the notion that vectors evaluated in this study did not result in off-target transgene expression. Consistent with previous reports that the reduction of GCAP1 in GC1KO mice is post-translational (i.e., mRNA levels are unchanged), we found no substantial changes in the levels of GCAP1 mRNA across samples (Fig. 5B).37

GNAT2 RNA in untreated GC1KO samples was reduced relative to GC1+/+ controls, a result likely due to the loss of cone photoreceptors in these retinas (Fig. 5B). In contrast, there were appreciable increases in GNAT2 mRNA levels in both AAV5- and AAV8(Y733F)-treated eyes, a result that further supports that cone photoreceptors are preserved in AAV-mGC1–treated GC1KO mice (Fig. 5B). Levels of rod PDE6α were relatively unchanged across samples, likely because rod photoreceptors do not degenerate in the GC1KO mouse (Fig. 5B).

**DISCUSSION**

Here we show that persistent AAV-mediated GC1 expression is capable of restoring retinal function and preserving cone photoreceptors in the GC1KO mouse over the long term (~1 year). This therapeutic longevity was characterized by a number of different criteria: the existence of GC1 protein in treated eyes at 15 months after treatment, the restoration of cone function as measured by electroretinography at 1 year after treatment, the increased cone survival in treated eyes at 11 months after treatment, and the recovery of vector genomes and GC1 mRNA in retinas at 1 year after treatment. When...
viewed as individual, discrete analyses, the sample sizes used in these assays were often small. However, when all are considered as correlates of therapeutic efficacy in mice exhibiting clear functional rescue (ERG), the sample size was effectively much larger. Within this context, therefore, we would suggest therapy persists beyond the period during which different vector serotypes were statistically compared for ERG rescue. This is the first demonstration of long-term therapy in a model of GC1 deficiency (~1 year) and builds on our previous report that AAV-mediated delivery of murine GC1 restored cone-mediated function and behavior and preserved cone photoreceptors in this model for 3 months.26

Cone photoreceptor function was significantly improved over the long term after AAV-mGC1 treatment, with AAV8(Y733F) vector being the most efficient. This result complements our previous report that an AAV8(Y733F) vector stably restored retinal structural and functional to the rd10 mouse, a model refractory to treatment with standard AAV vectors.36 Quantifying differences in rod amplitudes between treated and untreated eyes in the GC1KO mouse is complicated by the fact that rod function in this model is partially subserved by GC2 and by the large inter-animal variation in rod responses.38 Some improvements in rod-mediated responses in AAV8(Y733F)-treated GC1KO mice were observed more consistently than those recorded from AAV5-treated mice; however, these improvements did not reach statistical significance. This suggests that more aggressive expression of GC1 in the GC1KO eye can supplement the partial effect of GC2 on murine rod function.

Long-term preservation of cone photoreceptors was demonstrated 11 months after gene replacement. This result suggests that cone preservation is achievable over the lifetime of the animal. Although the preserved cones in treated GC1KO retina were not examined on an ultrastructural level (e.g., electron microscopy), our observation that cones remained functional over time by ERG analysis suggests that they were structurally intact. Long-term preservation of cone photoreceptors mediated by therapeutic AAV-GC1 has obvious clinical relevance because it suggests the potential to preserve macular cones and to restore usable daytime/color vision to patients with GC1 deficiency.

AAV-mediated GC1 expression persisted for at least 15 months after treatment (the latest time point evaluated by immunohistochemistry). Regardless of the serotype used or whether a photoreceptor-specific (hGRK1) or ubiquitous (smCBA) promoter drove its expression, GC1 was located exclusively in photoreceptors. Although transgene expression was limited to the target cell type, it should be noted that the hGRK1 promoter resulted in GC1 localization mainly within the proper compartment of the target cell (photoreceptor outer segments). This result, along with other successful proof-of-concept studies using this promoter, suggests that the hGRK1 promoter should be considered in the design of a clinical AAV vector targeting photoreceptors.39,40 However, because of the existence of multiple GRK isoforms across species that could limit the activity of hGRK1 in photoreceptor subtypes41 and instances in which promoter activity in the mouse retina does not translate perfectly to the nonhuman primate (NHP) retina,42 the activity of this promoter first must be thoroughly evaluated in the NHP retina.

Some evidence of retinal thinning was observed in two GC1KO retinal sections at 10 months after treatment with AAV5-smCBA-mGC1 or AAV5-hGRK1-mGC1. However, no appreciable thinning was observed in an older mouse that was evaluated 15 months after treatment with AAV5-hGRK1-mGC1. The small number of retinas analyzed precludes determination of the cause of thinning (injection-related damage or vector/transgene-related toxicity). Clearly, dose de-escalation experi-
ments in larger cohorts of mice will be required to determine whether retinal thickness can be preserved at lower therapeutic doses. This will be an essential experiment for firmly establishing any dose-limiting toxicity, a critical parameter in designing a safe dosing schedule for a clinical trial.

Despite the photoreceptor-exclusive nature of AAV-mediated GC1 expression, vector genomes were detected in the optic nerves and, in one instance, the brains of treated GC1KO mice. Our biodistribution data were collected from “diseased” animals. This is important because of evidence that patterns of vector transduction are different in diseased versus healthy retina and that this difference was due in part to vector distribution being enhanced in diseased retina. This is the first evaluation of biodistribution for an AAV vector containing a capsid surface–exposed tyrosine mutation. Although AAV5 is known to be ineffective for transducing ganglion cells of the mouse retina, some exposure of vector to retinal ganglion cells is expected as the syringe transverses the inner retina during subretinal injection and because the ratio of injection volume to total eye size is high in mice. The higher number of vector genomes detected in optic nerves of AAV8(Y733F)-treated eyes, therefore, could be due to the increased affinity of AAV8(Y733F) for retinal ganglion cells. At only one time point did we detect AAV5-delivered vector genomes in the brain of a treated GC1KO mouse with recovery only from the hemisphere opposite the injected eye. Our results (only 1 of 16 samples positive) are consistent with a previous study that reported a lack of AAV5-delivered sequence in brains of subretinally injected rats and dogs. By 10 months after injection, no vector genomes were recovered from brains of AAV5-treated GC1KO mice or from brains of mice treated with AAV8(Y733F) at any time point. Vectors used in this study were delivered at full strength. Future ex-
perimetric evaluation using cone-depleted retinas may show that a lower vector concentration is sufficient to affect therapy without spread to the brain. In general, other studies evaluating AAV biodistribution suggest that there is a progressive loss of vector sequences with time distant from the site of administration.\(^{29,47}\) Studies in rat, dog, and primate show that ocularly delivered AAV (serotypes 2, 4, and 5) do not spread widely outside the injected eye, with optic nerve and brain the most common exceptions.\(^{29,46,47}\) The lack of genome recovery from brains of treated GC1KO mice suggests that AAV5 and AAV8(Y733F) vectors used in this study have biodistribution patterns consistent with those reported for AVV2 by Jacobson et al.\(^{47}\)

Despite recovering vector genomes from optic nerves of treated eyes, immunostaining revealed a lack of GC1 expression in optic nerves of eyes treated with either AVV5-smCBA-mGC1 or AAV8(Y733F)-hGRK1-mGC1 vectors. A previous study by Stieger et al.\(^{48}\) detected transgene expression in optic nerves and brains of rats and dogs at 2 months and 4 weeks after subretinal injection with AAV8 containing green fluorescent protein (GFP). Taking into account that our AAV8(Y733F) vector contained the photoreceptor-specific hGRK1 promoter and our previous finding that GC1 expression was limited to photoreceptors even when under the control of a ubiquitous promoter such as smCBA, a lack of GC1 expression in optic nerves is not unexpected. Stieger et al.\(^{48}\) incorporated the strong, ubiquitous cytomegalovirus (CMV) promoter into their vector to drive GFP, a protein capable of being stably expressed in a wide variety of tissues when delivered by viral vectors.

Although both AVV5 and AAV8(Y733F) vectors were capable of providing long-term therapy to the GC1KO mouse, there are apparent advantages associated with using AAV8(Y733F). First and foremost, AAV8(Y733F) with a photoreceptor-specific promoter conferred significantly higher cone ERG responses to treated mice than either AVV5 vector. The reason for this may be the ability of AAV8 vectors to transduce areas outside the injection bleb in the rodent retina, whereas the area of retina transduced by AVV5 remains largely confined to the bleb.\(^{48}\) Thus AAV8(Y733F) may simply transduce, on average, a larger area of retina relative to AVV5. Although we did not evaluate the kinetics of vector-mediated GC1 expression in cones transduced with AAV8(Y733F) or AVV5 vectors in this study, it is known that AAV8(Y733F) supports a much faster onset of transgene expression than AVV5 vectors and that this temporal difference is important in other retinal degenerative models in terms of ability to preserve function.\(^{36,45}\) In the GC1KO mouse, the earlier availability of GC1 in AAV8(Y733F)-treated mice might have led to a higher overall number of preserved cone photoreceptors than that achieved in the AVV5-treated mice and, therefore, to more robust cone function. Again, experiments in NHP comparing the two serotypes are needed before firm conclusions can be drawn regarding which vector should be considered in a clinical setting.

This study provides insights into the effects of GC1 delivery to a mammalian model of GC1 deficiency. However, evaluating a gene replacement strategy for LCA1 in the GC1KO mouse has limitations. Rod photoreceptor structure and function are relatively preserved in this mouse, whereas LCA1 is characterized by extinguished cone and rod electroretinograms and by the degeneration of both cones and rods.\(^{23}\) With the goal to treat photoreceptors in an affected LCA1 retina, therapeutic vector transduction profiles should first be evaluated in a model that is more degenerative (i.e., exhibits more than just a loss of cone photoreceptors). Studies are under way to characterize these vectors in the recently described GC1/GC2 double knockout (GCdko) mouse, which exhibits loss of rod/cone function and both rod and cone degeneration.\(^{40,41}\) These studies should provide critical information regarding the recovery of rod function and whether, when treated together, both subclasses of photoreceptors are capable of being preserved.

The results presented here support further development of an AAV-based gene replacement therapy for LCA1 and potentially for GC1-related autosomal recessive CORD patients. Interestingly, our results may suggest that LCA12 patients could benefit from gene replacement as well. A recent report\(^{49}\) has shown that rd3, a gene associated with autosomal recessive photoreceptor degeneration in the rd3 mouse, rcld2 dog, and LCA12 patients, codes for RD3, a protein required for the export of GC1 from the endoplasmic reticulum to endosomal vesicles.\(^{50}\) Specifically, photoreceptor degeneration in the rd3 mouse is caused by impaired RD3-mediated guanylate cyclase expression and trafficking and is similar to the pattern exhibited by the GGrk1 mouse.\(^{49}\) Given the parallels in biochemical dysfunction, our results showing AAV-GC1 arrests cone degeneration and restores visual function would suggest that rd3 gene replacement may also be successful in the rd3 mouse and the rcld2 dog. Future gene therapy experiments using AAV-GC1 to treat GCdko mice will perhaps lend further support to this concept.

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