Gene Therapy Preserves Retinal Structure and Function in a Mouse Model of NMNAT1-Associated Retinal Degeneration

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INTRODUCTION
Nicotinamide mononucleotide adenylyltransferase 1 (NMNAT1)-associated retinal degeneration is an early-onset, recessive disease that causes severe vision loss during the first or second decade of life.1-5 The affected gene, NMNAT1, encodes a ubiquitously expressed enzyme that is essential for regenerating nicotinamide adenine dinucleotide (NAD)+ metabolism in tissues throughout the body. NMNAT1-associated disease is isolated to the retina. Since this condition is recessive, supplementing the retina with a normal copy of NMNAT1 should protect vulnerable cells from disease progression. We tested this hypothesis in a mouse model that harbors the p.Val9Met mutation in Nmnat1 and consequently develops a retinal degenerative phenotype that recapitulates key features of the human disease. Gene augmentation therapy, delivered by subretinal injection of adeno-associated virus (AAV) carrying a normal human copy of NMNAT1, rescued retinal structure and function. Due to the early-onset profile of the phenotype, a rapidly activating self-complementary AAV was required to initiate transgene expression during the narrow therapeutic window. These data represent the first proof of concept for a therapy to treat patients with NMNAT1-associated disease.

No treatment is available for nicotinamide mononucleotide adenylyltransferase 1 (NMNAT1)-associated retinal degeneration, an inherited disease that leads to severe vision loss early in life. Although the causative gene, NMNAT1, plays an essential role in nuclear nicotinamide adenine dinucleotide (NAD)+ metabolism in tissues throughout the body, NMNAT1-associated disease is isolated to the retina. Since this condition is recessive, supplementing the retina with a normal copy of NMNAT1 should protect vulnerable cells from disease progression. We tested this hypothesis in a mouse model that harbors the p.Val9Met mutation in Nmnat1 and consequently develops a retinal degenerative phenotype that recapitulates key features of the human disease. Gene augmentation therapy, delivered by subretinal injection of adeno-associated virus (AAV) carrying a normal human copy of NMNAT1, rescued retinal structure and function. Due to the early-onset profile of the phenotype, a rapidly activating self-complementary AAV was required to initiate transgene expression during the narrow therapeutic window. These data represent the first proof of concept for a therapy to treat patients with NMNAT1-associated disease.

Currently, no treatment exists for NMNAT1-associated retinal degeneration. Because patients incur considerable vision loss during the first years of life, but are expected to have normal longevity, an early intervention has the potential to preserve sight for many decades. Since this disease has a recessive inheritance pattern10 and because NMNAT1 is a relatively small gene,16 treatment with adeno-associated virus (AAV)-mediated gene augmentation therapy is an attractive strategy. At 840 bp, human NMNAT1 cDNA is well within the ~4.7-kb cargo capacity for single-stranded AAV (SS.AAV)17 and the ~2.2-kb cargo capacity of self-complementary AAV (SC.AAV).18 AAV-mediated gene augmentation is presently being used as a US Food and Drug Administration (FDA)-approved therapy for RPE65-associated retinal degeneration19 and in ongoing clinical20-23 and pre-clinical24-27 trials for other inherited retinal degenerations. In addition, an SC.AAV−SMN1 vector is approved for the treatment of spinal motor atrophy.28

When NMNAT1 was reported as a disease gene in 2012,1-4 a suitable animal model was not available for evaluating potential therapies in situ. Traditional Nmnat1 knockout mice were not viable,12 and conditional knockout animals, which have Nmnat1 ablated in targeted
retinal cells, would not have accurately represented the disease physiology. Subsequently, we identified and characterized an NMNAT1-associated retinal disease mouse model that is homozygous for the p.Val9Met (V9M) mutation in Nmnat1,14 an allele that has been found to cause retinal disease in members of unrelated families.4,10 Nmnat1V9M/V9M mice invariably develop an early-onset isolated retinal disease without obvious detriments to longevity or mobility, much like the humans they model. These mice have fully mature retinas and reliable responses to light at 3 weeks of age, as detected by electroretinogram (ERG), but a week later the photoreceptor layer shows signs of degeneration accompanied by reduced function.14 When the mice reach approximately 4 months of age, the retina is severely degenerated and responses to light stimuli are often undetectable.14

For the purpose of developing a therapy that preserves vision in people with NMNAT1-associated retinal degeneration, we used the Nmnat1V9M/V9M mouse model to test the hypothesis that the structure and function of the retina can be rescued if supplemented with normal NMNAT1. A human NMNAT1 cDNA was delivered to the retinas of Nmnat1V9M/V9M mice via recombinant AAV vectors that were evaluated independently. Efficacy varied across viral preparations and experimental conditions; therefore, we aimed to understand why specific variables were associated with success or failure and how these lessons might generalize to assist in the development of other AAV-mediated gene therapies.

RESULTS
DNA Construct and AAV Vector Preparation
A codon-optimized human NMNAT1 cDNA (Figure 1A) was incorporated into constructs that were then packaged into recombinant AAV vectors. Codon optimization has been reported to improve the level and duration of expression of human genes in transduced cells without altering the amino acid sequence of the protein product.29–33 All 174 nt substitutions introduced into the 840-bp NMNAT1 cDNA were silent, defining the normal human protein sequence.
the ubiquitously expressing CASI promoter was packaged into both a self-complementary and single-stranded version of AAV2/9. The self-complementary vector was selected for testing because it activates gene expression more rapidly than traditional single-stranded vectors. A construct containing an EGFP (enhanced green fluorescent protein) reporter gene, also driven by the CASI promoter and followed by the woodchuck hepatitis virus posttranslational regulatory element (WPRE) that serves to enhance AAV-mediated transduction in mouse retina, was packaged into SS.AAV2/9. This EGFP vector was spiked in with the SC.AAV2/9 vector (Figure 1B) and SS.AAV2/9 vector (Figure 1C) at $1 \times 10^6$ genomic copies per microliter (gc/µL) just before delivery to the mice so that the injected region of the retina could be identified by in vivo and ex vivo imaging. Another construct was made in which NMNAT1 was driven by the ubiquitously expressing CAG promoter and followed by a T2A cleavage sequence, EGFP, and then WPRE. After translation in the cell, the NMNAT1-EGFP fusion protein was enzymatically separated at the T2A cleavage site to avoid disruption of nominal protein conformations and kinetics. The construct was packaged into AAV2/Anc80, a synthetic AAV vector that was generated by ancestral sequence reconstruction and that can transduce retinal cells efficiently both in mice and in non-human primates (Figure 1D). Finally, the same CASI.NMNAT1 construct as described above was packaged into the AAV2/7m8 vector, which has been reported to transduce all retinal layers in mice following intravitreal injection (Figure 1E). The SC.AAV2/9, SS.AAV2/9, and AAV2/Anc80 vectors were delivered by subretinal injection, whereas the AAV2/7m8 vector was delivered by intravitreal injection.

**Gene Augmentation Using the Self-Complementary Vector Preserves Retinal Structure**

Gene augmentation therapy using the SC.AAV2/9 vector stably preserved retinal structure in a dose-dependent manner when administered to 2-week-old Nmnat1<sup>V9M/V9M</sup> mice. This finding was determined by measuring photoreceptor layer thickness in images collected in vivo using optical coherence tomography (OCT). Measurements from injected eyes were taken near the injection site in the inferior retina where rescue was anticipated to be most robust and in the superior retina that was far from the injection site. Measurements in the non-injected fellow eyes of both treated Nmnat1<sup>V9M/V9M</sup> mice and treated age-matched wild-type control mice were acquired in the inferior retina in the plane equivalent to where the measurements were collected in the injected eyes (Figure 2).

Intermediate and high titer injections of the SC.AAV2/9 vector, $1 \times 10^8$ and $2 \times 10^8$ gc/µL, respectively, each provided significant rescue across the injected retina (Figure 3A). For example, in 9-month-old mutant mice, the average thicknesses of the inferior and superior photoreceptor layers of retinas injected with the $2 \times 10^8$ gc/µL dose were $100.0 \pm 4.8$ and $96.1 \pm 6.4$ µm, respectively, whereas measurements in the inferior region of the fellow non-injected eyes averaged $31.3 \pm 0.8$ µm (68.7 µm difference from injected inferior retina, $p < 0.0001$, and 64.8 µm difference from superior injected retina, $p < 0.0001$). Photoreceptor layer thickness in treated retinas of mutant mice were typically within ~20% of those collected from non-injected inferior retinas of wild-type mice. Within the treated mutant retinas, values tended to be similar ($p > 0.05$) at the inferior (near the injection site) and superior (far from the injection site) locations. However, photoreceptor layer thickness decreased by ~20% with distance from the injection site following the $1 \times 10^8$ gc/µL dose. Structural rescue following treatment with the lowest titer tested for this vector, $1 \times 10^7$ gc/µL, was more modest.

Because the impairment of NMNAT1 activity may differ by mutation, understanding whether overexpression of this enzyme following gene augmentation, the viral particles, or the viral titer can cause toxicity is important. For this purpose, wild-type mice were injected with the SC.AAV2/9 vector using the three abovementioned titers. Across data collection time points for all doses, photoreceptor layer thickness was typically unaffected, with small but statistically significant differences being noted in only three instances (Figure 3B). The largest disparity was in the superior region of the injected retinas in which the thickness was decreased by $17.9 \pm 6.9$ µm ($p = 0.039$) 9 months
after the $2 \times 10^9$ gc/μL injection. However, given that the inferior region (near the injection site) of the same retina was unaffected, this decrease was unlikely to be due to toxicity.

Single-Stranded Vectors Do Not Stably Preserve Retinal Structure

In contrast to the SC.AAV2/9 vector, the SS.AAV2/9, AAV2/7m8, and AAV2/Anc80 vectors did not provide stable structural rescue of the Nmnat1V9M/V9M retina. At 2 months of age, the inferior region of mutant retinas injected with the SC.AAV2/9 vector showed a modest rescue of the photoreceptor layer thickness ($11.4 \pm 4.3$ μm, $p = 0.033$) versus the same region of the non-injected fellow retina (Figure 3C), whereas the AAV2/7m8 vector had no effect (Figure 3D). The $5.5 \times 10^8$ gc/μL dose of the AAV2/Anc80 vector produced transient rescue in the mutant retina that did not persist beyond 2 months of age and was confined to the inferior region. At the 1.5- and
To test whether the dilution buffer used with the untreated fellow retinas (Figure 3E). No rescue was observed when the dose of the AAV2/Anc80 vector was decreased to 1 × 10^8 gc/μL (data not shown), and increasing the dose to ≥1 × 10^9 gc/μL failed to apparent toxicity. The high titer injections of the AAV2/Anc80 vector caused detachment in some retina. (G) A montage of the full retinal section, labeled with the EGFP expression (green). (D–F) Both signals are detected in the injected fellow retina. (A–C) The non-injected retina has neither α-NMNAT1 antibody reactivity (red) nor EGFP expression (green). (D–F) Both signals are detected in the injected fellow retina. (A–C) The non-injected retina has neither α-NMNAT1 antibody reactivity (red) nor EGFP expression (green). (D–F) Both signals are detected in the injected fellow retina. (A–C) The non-injected retina has neither α-NMNAT1 antibody reactivity (red) nor EGFP expression (green). (D–F) Both signals are detected in the injected fellow retina.

2-month time points, respectively, the inferior region of AAV2/ Anc80-treated retinas had photoreceptor thicknesses that were 42.0 ± 3.7 μm (p = 0.0005) and 27.5 ± 9.9 μm (p = 0.045) greater than that of the untreated fellow retinas (Figure 3E). No rescue was observed when the dose of the AAV2/Anc80 vector was decreased to 1 × 10^8 gc/μL (data not shown), and increasing the dose to ≥1 × 10^9 gc/μL failed to apparent toxicity. The high titer injections of the AAV2/Anc80 vector caused detachment in some retina. (G) A montage of the full retinal section, labeled with the EGFP expression (green). (D–F) Both signals are detected in the injected fellow retina. (A–C) The non-injected retina has neither α-NMNAT1 antibody reactivity (red) nor EGFP expression (green). (D–F) Both signals are detected in the injected fellow retina. (A–C) The non-injected retina has neither α-NMNAT1 antibody reactivity (red) nor EGFP expression (green). (D–F) Both signals are detected in the injected fellow retina.

Non-injected Injected (fellow)

Figure 4. Ex vivo Imaging Shows Structural Rescue at Cellular Level in Retina Treated with the SC.AAV2/9 Vector

Immunolabeled retina from a representative 9-month-old Nmnat1^V9M/V9M mouse. (A–C) The non-injected retina has neither α-NMNAT1 antibody reactivity (red) nor EGFP expression (green). (D–F) Both signals are detected in the injected fellow retina. (G) A montage of the full retinal section, labeled with the α-NMNAT1 antibody, shows the extent of transgene expression across the retina. DAPI (blue) is the counterstain in (C), (F), and (G), and the white arrowhead indicates the injection site in (G). Images shown are from a 9-month-old Nmnat1^V9M/V9M mouse injected on P14 with SC.AAV2/9 at a titer of 2 × 10^9 gc/μL. Original magnification, ×63; scale bar represents 50 μm. RPE, retinal pigment epithelium; OS, outer segments; ONL, outer nuclear layer; INL, inner nuclear layer; GCL, ganglion cell layer.

Nuclear Localization of Human NMNAT1 is Observed across Cell Types in SC.AAV2/9-Treated Retinas

Ex vivo immunofluorescence imaging of retinas treated with the SC.AAV2/9 vector was performed using a custom chicken polyclonal α-human NMNAT1 antibody. This antibody displays no cross-reactivity with mouse NMNAT1 (Figure S2). Treated retinas showed nuclear localization of human NMNAT1 in the retinal pigment epithelium (RPE), outer nuclear layer, inner nuclear layer, and ganglion cell layer, indicating transgene expression across cell types (Figures 4D–4F). Furthermore, an injection can transduce more than two-thirds of the tissue area (Figure 4G).

Gene Augmentation Using a Self-Complementary Vector Preserves Retinal Function

To assess whether structural rescue of the retina translated to preservation of function, we measured rod and cone photoreceptor function in vivo with ERG testing, a non-invasive procedure that records electrical responses of the retina to light stimulation. The intensity of the light was calibrated so that rod-mediated responses and mixed rod/cone-mediated responses were recorded separately when the mice were dark-adapted, and cone-mediated responses were recorded when the mice were light-adapted. For each stimulus condition, the magnitude of the ERG b-wave, which is mediated by ON bipolar cells, was used as an indirect indicator of the photoreceptor response and a measure of inner retinal function. To directly assess photoreceptor function, the ERG a-wave was analyzed for the mixed rod/cone response; this near maximum stimulus condition was selected on the basis that, in normal mice, the a-wave is more robust in mixed rod/cone ERGs than in dimmer stimulus-elicited dark-adapted ERGs or in light-adapted ERGs.

Retinal function was preserved in eyes of mutant mice treated with the SC.AAV2/9 vector. With injection of the 2 × 10^9 gc/μL dose, the b-wave was significantly greater than that of the untreated fellow eye by 4 months of age for each stimulus condition and remained so through 6 months of age for the light-adapted condition. The b-wave remained significantly greater to at least 9 months of age for both dark-adapted conditions, and cone-isolated ERGs recorded at 9 months of age tended to be greater in the treated mutant eye than in the untreated fellow eye (Figure 5A, top row). For example, in 6-month-old mutant mice, the b-wave of the treated retina exceeded that of the fellow untreated retina for each condition: 156.1 ± 23.4 versus 36.0 ± 7.1 μV (p = 0.0013) for the rod response, 293.6 ± 51.1 versus 62.0 ± 11.8 μV (p = 0.0034) for the mixed rod/cone response, and 101.3 ± 18.6 versus 25.3 ± 6.2 μV (p = 0.0067) for the cone response. Likewise, the a-wave was significantly greater for the treated eyes of the mutant mice than for the fellow untreated eyes by 4 months of age, and this effect remained through 9 months of age (Figure S3).

The responses associated with rescue in the Nmnat1^V9M/V9M retinas were consistently lower than those of the untreated wild-type retinas, and vector titer may have contributed to this effect. ERGs recorded from injected eyes of 9-month-old wild-type mice tended to be negatively correlated with dosage; the average response amplitude across the three stimulation conditions was 86% of normal with use of the 1 × 10^8 gc/μL titer versus 73% of normal with the 2 × 10^9 gc/μL titer. Regardless, the structure and implicit times of the rescued ERG waveforms were normal at all ages (Figure 5A, bottom row). The gradual
decline in ERG amplitudes observed in wild-type animals between 2 and 9 months of age is normal.44,51

The SC.AAV2/9 vector at a lower dose of $1 \times 10^8$ gc/µL still showed evidence of functional rescue in the injected mutant retina (Figure 5B), whereas no sign of functional rescue by the AAV2/Anc80 vector at a similar dose of $5.5 \times 10^8$ gc/µL was observed at 2 months of age (Figure 5C), despite evidence of preserved retinal structure.

Self-Complementary Vector Activates NMNAT1 Expression Earlier Than Single-Stranded Vectors

To understand why retinal structure and function were best preserved when treatment was delivered via the SC.AAV2/9 vector, we compared the early NMNAT1 expression profiles of SCAAV2/9, SS.AAV2/9, and AAV2/Anc80 delivery vectors in treated eyes of mixed C57BL/6J-129S6 wild-type mice. Injections occurred at around postnatal day 16 (P16), and each vector was used with a titer of $1 \times 10^8$ gc/µL. At 14 days post-injection, timed precisely for each animal, immunolabeling of NMNAT1 delivered by the SC.AAV2/9 vector was observed in all retinal layers with particularly high density in the outer nuclear layer (Figure 6A). Conversely, in SS.AAV2/9- and AAV2/Anc80-injected retinas, detection of NMNAT1 was sparse with relatively few photoreceptors labeled, even though the SS.AAV2/9 vector carried identical cargo and the AAV2/Anc80 vector has been shown to transduce the mouse retina photoreceptors with wide coverage.37 NMNAT1 was undetectable at 7 days post-injection, regardless of which of the three delivery vectors was used (data not shown). The greater efficiency of rod photoreceptor transduction by AAV2/Anc80 that we have reported previously37 may be due to those mice having been injected at substantially older ages (6–8 weeks old) than the mice described herein.

A Successful Intervention Requires Transduction of Photoreceptors

To determine whether an earlier intervention would produce better efficacy from the AAV2/Anc80 vector, injections were performed in neonatal mice at P0 through P2. While most cell types, including cone photoreceptors, showed strong NMNAT1 expression in these animals, expression in rod photoreceptors was relatively weak (Figure 6B); cones were distinguished by the location of their nuclei in the outermost rows of the outer nuclear layer (ONL).37 Given that AAV2/Anc80 efficiently transduces rod photoreceptors following subretinal injection in adult mice (6–8 weeks old),37 and since another research group reported poor transduction of rods by other AAV vectors following subretinal injection of neonatal mice,37 we did not pursue neonatal injections using the SS.AAV2/9 or SC.AAV2/9 vectors. Similar to AAV2/Anc80 injection of neonatal mice, intravitreal delivery of NMNAT1 via AAV2/7m8 in 2-week-old mice did not preserve the photoreceptor layer, as observed by OCT 4 weeks after injection, and we determined by immunohistochemistry that NMNAT1 expression was absent in the outer retina, despite a strong inner retina signal (Figure 6C).

DISCUSSION

This study is the first demonstration of a gene therapy that successfully treats NMNAT1-associated retinal degeneration. Retinas of Nmnat1<sup>V9M/V9M</sup> mice supplemented with a normal copy of human NMNAT1 via AAV-mediated gene augmentation show preservation of both structure and function. This result was achieved using a self-complementary AAV vector that was more effective than single-stranded vectors delivering the same transgene. Data indicate that the rapid activation of gene expression associated with the self-complementary vector provided a better outcome by permitting widespread transduction of retinal cells within the narrow therapeutic window for this disease. Efficient transduction of photoreceptor cells appears to be necessary for therapeutic efficacy, and rescue of retinal structure tended to be more robust than that of function. The results presented suggest that, with further optimization, AAV-mediated gene augmentation therapy has the potential to benefit patients affected by NMNAT1-associated retinal degeneration.

Retinas of Nmnat1<sup>V9M/V9M</sup> mice supplemented with a normal copy of human NMNAT1 via AAV-mediated gene augmentation show preservation of structure and function for at least 9 months. This effect, which was dose-dependent, was achieved using the SC.AAV2/9 delivery vector. Although the highest viral titer used conferred the greatest effect, the observation that an intermediate titer also provided rescue suggests that high doses of AAV-NMNAT1 may not be needed for a clinically effective therapy. In contrast to the SC.AAV2/9 vector, the three single-stranded AAV vectors did not provide sustained therapeutic effects. The SS.AAV2/9 vector, the direct counterpart of the SC.AAV2/9 vector, afforded only modest preservation of retinal structure in mice at 2 months of age. Similarly, the AAV2/Anc80 vector provided a transient structural benefit with no functional rescue. The AAV2/7m8 vector, delivered by intravitreal injection, did not transduce the outer retina and failed to provide a structural or functional benefit. In comparison to these single-stranded vectors, the self-complementary vector had a transgene expression profile that was comparatively faster, greater, and included more target cells, notably photoreceptors.

Figure 5. ERG Shows Preservation of Retinal Function after Treatment with SC.AAV2/9 Vector

(A) Retinas from Nmnat1<sup>V9M/V9M</sup> mice treated with a $2 \times 10^8$ gc/µL dose of SC.AAV2/9 generated significantly larger rod, mixed rod/cone, and cone-isolating ERGs than did the untreated fellow retinas, as measured by the ERG b-wave. Measurements from the non-injected Nmnat1<sup>V9M/V9M</sup> retinas are compared to injected fellow retinas and to non-injected and injected retinas of wild-type mice (age 2, 4, 6, 9 months: n = 6, 4, 14, 13 wild-type; 5, 9, 10, 9 mutant; top row). ERG waveforms from a 6-month-old treated Nmnat1<sup>V9M/V9M</sup> mouse (treated retina, blue trace; untreated retina, red trace) and an age-matched wild-type littermate (untreated retina, black trace; bottom row). (B) ERG measurements to 9 months of age for mice injected with SC.AAV2/9 vector at the $1 \times 10^8$ gc/µL dose (age 4, 6, 9 months: n = 5, 3, 3 wild-type; n = 5, 4, 4 mutant). (C) ERG measurements to 4 months of age with the AAV2/Anc80 vector at the $5.5 \times 10^8$ gc/µL dose (age 2, 4 months: n = 1, 5 wild-type; n = 9, 5 mutant). Error bars represent the SEM. *p < 0.05, **p < 0.1, ***p < 0.001, ****p < 0.0001.
The rapid activation of gene expression associated with the self-complementary vector permitted efficient transduction of rod photoreceptors within the narrow therapeutic window for this disease, leading to a substantially better outcome than for single-stranded vectors. The transgene is expressed quickly due to the double-stranded configuration of the self-complementary genome, which allows the rate-limiting step of second-strand synthesis to be circumvented. The Nmnat1<sup>Y8M/Y8M</sup> mouse model has a narrow therapeutic window, bounded on the early side by immature rods that are unable to be transduced efficiently by AAV<sup>SS</sup> and on the late side by the onset of the rapidly progressing disease. We hypothesize that the shorter latency for gene expression after transduction by the SC.AAV2/9 vector allowed for a more efficient treatment within these constraints.

Treatment with the self-complementary vector resulted in successful transduction of photoreceptors, which led to substantially better preservation of retinal structure and function. Two-week-old (i.e., ~P16) mice were injected because AAV has been reported to transduce approximately twice as many rods when it is delivered at P21 versus P10 and has also been observed to have poor transduction of rods when delivered at neonatal ages. The injections were not delayed further because, despite the rapid transgene activation, more than a week was required to detect NMNAT1 expression in retinas of wild-type mice dosed with the SC.AAV2/9 delivery vector. SC.AAV has been observed in previous studies to produce low-level transgene expression in retinal cells that is detectable by immunofluorescence microscopy by 7 days post-injection. In the present study, it was not determined whether the relatively young age of the mice at the time of injection, the viral vector, or the DNA construct may have contributed to the latency in transgene expression. Because the co-injected EGFP was delivered via the more slowly activating SS.AAV, this marker was not informative for evaluating whether the NMNAT1 transgene was rate-limiting. Furthermore, we found that early intervention with SS.AAVs was unsuccessful, likely because they did not transduce rods. Future experiments using cell type-specific promoters could help identify whether NMNAT1 supplementation in photoreceptors is sufficient for treatment efficacy, and, if not, which other cell types are required. Success of this vector may also be due to higher levels of gene expression, and this will be further examined in future studies, as well.

In eyes treated with the SC.AAV2/9 delivery vector, the level of preservation was consistently greater and more stable for retinal structure than for function. The photoreceptor layer was only ~15%–25% thinner in treated mutant retinas than in wild-type retinas; however, ERG b-wave and a-wave amplitudes tended to be ~50% lower in treated mutant retinas as compared to untreated wild-type retinas. Despite early preservation of retinal structure in treated eyes of the mutant mice, functional rescue did not become apparent by ERG until after the 2-month time point. While the mechanism underlying this latency will require further investigation, the transient lack of a-wave rescue indicates direct photoreceptor involvement. Furthermore, the photoreceptor layer of wild-type retinas injected with the SC.AAV2/9 vector at the 2 x 10<sup>5</sup> gc/µL dose was only slightly thinner (~1%–8%) than the same region of retina in the fellow non-injected eyes, but retinal function was reduced by ~25%. It is possible that the reduced ERG signal could be due to mechanical trauma from the injection procedure; however, damage to a small region would likely not affect the full-field ERG to this extent. Furthermore, ERG responses were not decreased in wild-type mice treated with the

**Figure 6. Vector-Specific Transgene Expression Profiles Indicate That Treatment of Photoreceptors Is Required for a Successful Therapy**

(A) SC.AAV2/9, SS.AAV2/9, and AAV2/Anc80 vectors were delivered at 1 x 10<sup>8</sup> gc/µL to wild-type mice at age ~P16. At precisely 14 days post-injection, dense α-NMNAT1 immunoreactivity (red) is visible in the retina (notably in the ONL) injected with the SC.AAV2/9 vector (left column), whereas this signal is sparse in age-matched retinas injected with either the SS.AAV2/9 (center column) or the AAV2/Anc80 (right column) vector. Original magnification, ×40; scale bar represents 75 µm. (B) A retina from a 5.5-week-old wild-type mouse that was injected with 3.5 x 10<sup>9</sup> gc/µL of AAV2/Anc80 at P1 shows strong α-NMNAT1 immunoreactivity (red) in virtually all cell types except for rod photoreceptors at 5 weeks post-injection. The labeled cells populating the outermost row of the ONL (flanked by green arrowheads) are cones. Original magnification, ×20; scale bar represents 100 µm. (C) Intravitreal injection of the AAV2/7m8 vector at 3 x 10<sup>9</sup> gc/µL in a 2-week-old control mouse generates strong NMNAT1 expression (red) in the INL and GCL but not in the ONL at 4 weeks post-treatment. DAPI (blue) is the counterstain in the bottom row of each panel. Original magnification, ×20; scale bar represents 100 µm. ONL, outer nuclear layer; INL, inner nuclear layer; GCL, ganglion cell layer.
$1 \times 10^6$ gc/μL titer of SC.AAV2/9, suggesting that the reduced retinal function observed following injection of the higher dose of SC.AAV2/9 could be due to some toxicity of the vector. High levels of NMNAT1 itself could be associated with reduced retinal function, although this seems unlikely based on results of previous studies showing that high levels of NMNAT1 and NAD⁺ are not detrimental to neural cells. If excessive NMNAT1 can be toxic to cells, then with high titer delivery of the AAV2/Anc80 vector, inclusion of WPRE in the DNA construct may have driven transgene expression to levels that cause an inflammatory response. Regardless, the potential for toxicity by NMNAT1 overexpression in the retina needs to be investigated further. Other possible sources of toxicity include elements of the viral vector or a contaminant in the vector preparations, although we found no specific evidence for these candidates. It is possible that the CASI promoter, similar to some other broadly active promoters, could contribute some toxicity to AAV vectors, but further work is needed to assess this possibility.

The therapeutic effects of NMNAT1 gene therapy reported in the present study provide evidence that gene augmentation therapy could be used to treat patients suffering from NMNAT1-associated retinal degeneration. Self-complementary AAV vectors have been successfully used to treat other neurodegenerative disorders such as spinal motor atrophy. As indicated, additional studies will be needed to optimize NMNAT1 gene therapy for clinical translation. For example, the ability of this therapy to target both cone and rod photoreceptors is likely to be important for clinical translation because foveal atrophy is a common feature associated with NMNAT1 disease. For patients with intact foveas, maintaining healthy rods would support the long-term viability of the treated central and peripheral cones, and for patients with foveal atrophy, treatment outside of the macula to preserve sight via peripheral cones and rods could also provide considerable benefit. Because gene augmentation has the potential to preserve vision in NMNAT1-disease patients from childhood onward, the availability of such a therapy would improve quality-of-life for this global patient population, especially in regions where blindness is associated with poverty, denial of access to education, and increased mortality.

MATERIALS AND METHODS

Mouse Lines

The pV9M-Nmnat1 mouse line was derived previously from an N-ethyl-N-nitrosourea (ENU) mutagenesis screen. To increase fecundity, the original C57BL/6J line was alternately outcrossed with wild-type 129S6/SvEvTac mice (Taconic, Rensselaer, NY, USA) and wild-type C57BL/6 mice (The Jackson Laboratory, Bar Harbor, ME, USA) to maintain a mixed C57BL/6J-129S6 genetic background. Wild-type CD1-IGS mice (Charles River, Wilmington, MA, USA), used only to screen for vector component toxicity, were maintained separately. Male and female mice were used in experiments without preference.

Animal Husbandry

Mice were fed 4% fat rodent diet and water ad libitum and housed in a 12-h light/12-h dark cycle. This study conformed to the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Vision Research, and all procedures were approved by the Animal Care and Use Committee.

Genotyping

A tissue biopsy was prepared for polymerase chain reaction (PCR) using All-In-One mouse tail direct lysis buffer (Allele Biotech, San Diego, CA, USA) according to the manufacturer’s instructions. PCR was performed using the forward primer 5′-CATG GCTGTGCTGAGGTG-3′ (intron 1) and reverse primer 5′-AACA GCCTGAGGTGCATTT-3′ (exon 2) to amplify a 691-bp region of Nmnat1 that includes codon 9. The 20-μL PCR reactions had final concentrations of 200 μmol/L for each primer, 200 nmol/L for each of the deoxyribonucleotide triphosphates (dNTPs) (deoxyadenosine triphosphate [dATP], deoxyguanosine triphosphate [dGTP], deoxythymidine triphosphate [dTTP], and deoxycytidine triphosphate [dCTP]), 2 mmol/L MgCl₂, and 1 U of Hot FirePol DNA polymerase (Solis BioDyne, Tartu, Estonia). The thermocycling protocol was 95°C for 14 minutes; 30 cycles of 95°C for 45 s, 53°C for 45 s, 72°C for 30 s; and 72°C for 7 min. Next, the amplified product was subjected to Sanger sequencing using primer 5′-ACGTATTGCCAACCTGTCTC-3′, and the resulting data were then analyzed at c.25 to identify each mouse as being wild-type, Nmnat1 WT/Y9M, or Nmnat1 V9M/V9M.

DNA Construct and AAV Vector Preparation

A codon-optimized human NMNAT1 cDNA (Figure 1A), designed by DNA 2.0 (Menlo Park, CA, USA) and synthesized into a gBlock gene fragment (Integrated DNA Technologies, Coralville, IA, USA), was incorporated into constructs that were then packaged into recombinant AAV viral vectors. Plasmids containing the complete constructs were generated using standard endotoxin-free molecular cloning techniques and validated by sequencing NMNAT1 and regions crossing ligation sites. AAV was prepared by the Grousebeck Gene Therapy Center of Massachusetts Eye and Ear, as described previously. Purified virus was collected and titered in a final buffer containing 1× phosphate-buffered saline (PBS), 35 mM NaCl, and 0.001% Pluronic F68 surfactant. The same buffer was used to further dilute the virus, if required, to achieve the target dose. To assist with the injection procedure, ~0.25% of fluorescein (AK-Fluor, Akorn, Lake Forest, IL, USA) was mixed into the working solution as a tracer.

Anesthesia

For general anesthesia, a mixture of ketamine/xylazine was delivered by intraperitoneal injection. Two-week-old mice received a dose of 37.5 mg/kg ketamine and 3.8 mg/kg xylazine and adult mice received a dose of 100 mg/kg ketamine and 20 mg/kg xylazine. To counteract the formation of anesthesia-induced corneal opacities, a 2 mg/kg dose of yohimbine HCl (Wedgewood Pharmacy, Swedesboro, NJ, USA) was administered by subcutaneous injection immediately after each recovery procedure in which ketamine/xylazine was used. For neonatal mice, general anesthesia by hypothermia was induced by indirect exposure to ice.
Virus Delivery
In 2-week-old mice, a Micro4 microinjection pump with an RPE kit (World Precision Instruments, Sarasota, FL, USA) was used to deliver the viral vectors into either the subretinal space or the vitreous chamber. Pupils were dilated using either tropicamide (1%) or a mixture of tropicamide (0.25%), phenylephrine HCl (1.25%), and cyclopentolate (0.5%). Mice were anesthetized with ketamine/xylazine, and local anesthesia was administered topically using proparacaine HCl (0.5%). Next, the eye was propoded and a 30G syringe needle was used to puncture the superior-temporal sclera and retina immediately posterior to the limbus to make an entry route for a blunt-end 33G cannula.

For subretinal injections, the cannula tip was positioned in the subretinal space of the inferior-nasal quadrant of the eye, 0.75 μL of vector was injected, and the formation of a bleb was confirmed by visualization. The cannula was held in place for approximately 3 s following injection to avoid reflux of the vector and then gently removed from the eye. Finally, the entry wound was treated by tamponade with a cotton swab. The eyes were then hydrated with artificial tears (Blink Tears, Abbott Laboratories, Chicago, IL, USA), and the mice recovered from anesthesia on a heating pad. The procedure for intravitreal injection of two-week-old mice was identical, except that the cannula tip was positioned in the center of the vitreous chamber during injection.

The FemtoJet 4i microinjection system (Eppendorf, Hamburg, Germany) was used to deliver subretinal injections to neonates. While the mice were anesthetized on ice, the tip of a 30G hypodermic needle was used to separate the upper and lower eyelids. The eye was propoded and a custom beveled glass needle (catalog no. C060609, Origio, Trumbull, CT, USA) was directly inserted through the sclera and positioned in the underlying subretinal space. A single bolus of 0.5 μL of vector was administered, after which the needle was held in place for approximately 3 s to avoid reflux and then gently removed. The mice recovered from anesthesia on a heating pad.

In Vivo Retinal Imaging
En face and cross-sectional images of the retina were acquired using fundus photography and spectral domain OCT, respectively, and as described previously14 with the following modifications: some fundus images were taken through a filter that allows for visualization of EGFP and therefore early evaluation of AAV injection quality, the rectangular volume OCT scans were taken at multiple locations across the retina so that inferior and superior regions could be measured accurately, and 10 B-scans were registered/averaged to generate final OCT images. Using InVivoVue OCT software (Bioptigen), four equally spaced electronic caliper measurements were made from the outer plexiform layer to the RPE to measure photoreceptor layer thickness. Across the time points tested (2, 4, 6, and 9 months, unless otherwise stated) for each reagent, the sample sizes were as follows: SC.AAV2/9 1 × 10^8 gc/μL (n = 8, 8, 5, wild-type; n = 8, 7, 5, 4 mutant), SC.AAV2/9 1 × 10^9 gc/μL (n = 8, 8, 4, 4 wild-type; n = 3, 4, 4, 4 mutant), SC.AAV2/9 2 × 10^8 gc/μL (n = 16, 16, 16, 15 wild-type; n = 10, 10, 9, 9 mutant), SS.AAV2/9 1 × 10^8 gc/μL (age 2 months: n = 10 wild-type; n = 9 mutant), AAV2/7m8 3 × 10^8 gc/μL (age 2 months: n = 3 wild-type; n = 3 mutant), AAV2/Anc80 5.5 × 10^8 gc/μL (age 1.5, 2, and 4 months, n = 4, 9, 13; wild-type, n = 4, 8, 15 mutant).

Electroretinography
Full-field, flash ERGs were collected from the mice as described previously. Briefly, mice were dark adapted overnight, and rod responses and mixed rod/cone responses were generated using a 0.01 cd·s/m² (scotopic) and 10 cd·s/m² (photopic) broadband light stimulus, respectively. Next, the mice were light adapted by exposure to a steady 30 cd/m² (photopic) broadband light for 10 min; this rod-suppressing light remained on in the background during the acquisition of cone-isolated responses to a 20 cd·s/m² (photopic) broadband light stimulus. Across the time points and stimulus conditions tested for each reagent, the sample sizes were as follows: SC.AAV2/9 1 × 10^8 gc/μL (age 4, 6, 9 months: n = 5, 3, 3 wild-type; n = 5, 4, 4 mutant), SC.AAV2/9 2 × 10^8 gc/μL (age 2, 4, 6, 9 months: n = 6, 4, 14, 13 wild-type; 5, 9, 10, 9 mutant), AAV2/Anc80 5.5 × 10^8 gc/μL (age 2, 4 months: n = 1, 5 wild-type; n = 9, 5 mutant).

Statistical Analysis
Statistical analyses were completed in Prism version 8.2.1 (GraphPad, San Diego, CA, USA). For OCT and ERG time courses, a two-way ANOVA was performed using a mixed-effects regression model. When analyzing the effect of a treatment, the inferior retina of the non-injected eye was used as the negative control to which the means of all other measurements were compared. The Dunnett post hoc test was used to account for type I error generated from multiple comparisons testing. Analyses of the effects of injection and all quantitative data are reported as the mean ± SEM.

Anti-Human NMNAT1 Antibody Development
Purified, full-length human NMNAT1 protein was used by Aves Labs (Tigard, OR, USA) as the antigen to generate polyclonal antibodies in chicken. The resulting antibody reacts strongly against human NMNAT1 without cross-reacting with the murine ortholog (Figure S2).

Immunohistochemistry and Ex Vivo Retinal Imaging
Mice were euthanized by CO2 asphyxiation and immediately perfused through the heart using a Masterflex peristaltic pump (Cole-Parmer, Vernon Hills, IL, USA). Each animal was perfused first with 0.13 mol/L PBS (pH 7.2–7.4) that contained 2 U of heparin/mL until the perfusate became clear, and this was followed by perfusion of ~40 mL of 2% paraformaldehyde (PFA). Both solutions were warmed to ~37°C at the time of perfusion. A small vessel cautizer (catalog no. 180000-00, Fine Science Tools, Foster City, CA, USA) was used to mark the cornea immediately anterior to the superior limbus. Eyes were enucleated, incubated at 2% PFA for 30 min at room temperature, the anterior segment was removed, and then the remaining eye cup was incubated once again in 2% PFA for 30 min at room temperature before being immersed in 30% sucrose at room temperature for

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at least 1 h. Eye cups were embedded, sectioned at 10-μM thickness by cryotomy, immunolabeled and stained, and imaged using either the Eclipse Ti fluorescence microscope (Nikon, Tokyo, Japan) or the TCS SP8 confocal microscope (Leica, Wetzlar, Germany), as described previously.1 The number of mice evaluated in Figure 6 were n = 2 per condition for (A), n > 4 for (B), and n = 3 for (C).

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.omtm.2020.07.003.

AUTHOR CONTRIBUTIONS
Conceptualization, E.A.P., L.H.V., and S.H.G.; Methodology, E.A.P., L.H.V., S.H.G., and R.X.; Formal Analysis, S.H.G., R.F., E.H., and M.J.S.; Investigation, S.H.G., R.F., E.H., M.J.S., E.E.B., and B.S.P.; Writing, S.H.G., E.A.P., and E.E.B.

CONFLICTS OF INTEREST
L.H.V. holds founder equity in GenSight Biologics, is a consultant to a number of biotech and pharmaceutical companies, and is an inventor on gene therapy patents, including Anc80, which are licensed to various entities. The remaining authors declare no competing interests.

ACKNOWLEDGMENTS
This research was supported by funding from NEI grant EY012910 (to E.A.P.), the Foundation Fighting Blindness and the Gavin R. Stevens Foundation (to E.A.P. and L.H.V.), the Sara Elizabeth O’Brien Trust (to S.H.G.), Fight for Sight (to S.H.G.), and NIH grant P30 EY014104 (to the MEEI Ophthalmology Core). Furthermore, the authors are grateful to Dr. Mathias Ziegler (University of Bergen) for providing purified human NMNAT1 that was used as the antigen for the antibody generation, and Dr. Rosario Fernandez-Godino (Massachusetts Eye & Ear) for providing cultured ARPE-19 cells.

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