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**SMALL ET AL.: 'Carolina macular dystrophy is caused by dysregulation of the retinal transcription factor**

**PRDM13.' OPTHALMOLOGY vol. 123, no. 1, October 2015, pages 9 - 18, XP029364142**

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# DESCRIPTION

## BACKGROUND OF THE DISCLOSURE

**[0001]** The present disclosure relates to using CRISPR-based methods to perform gene editing in patients in order to treat autosomal dominant diseases.

**[0002]** There are currently no cures for numerous autosomal dominant or recessive diseases that have a profoundly negative impact on quality of life. Dominant forms of retinitis pigmentosa (adRP), cone-rod dystrophies and juvenile macular degenerations are prime examples of dominant autosomal diseases that affect the eye. These autosomal dominant diseases are characterized by the presence of a mutant gene expressing a defective protein. These diseases are not, therefore, readily amenable to therapies that simply add a normal, healthy gene (so called "gene supplementation" or "gene addition"), since the disease causing gene and protein are still present. Instead, gene editing offers the only means to directly repair the defective gene and, thus, the most promising therapeutic strategy.

**[0003]** Retinal degenerative diseases affect at least 9 million Americans (Friedman DS et al. Arch Ophthalmol. 2004 Apr;122(4):564-72; Schmier JK et al. Pharmacoeconomics. 2006;24(4):319-34). Among the most devastating retinal degenerative disease is retinitis pigmentosa (RP), a common form of inherited neurodegeneration, which affects 1.5 million people worldwide and for which treatment is inadequate. RP is a degenerative eye disease that results in retinal degeneration and vision loss. Hereditary mutations in the rhodopsin gene (*RHO*) are the most common cause of autosomal dominant RP, accounting for 20-30% of the cases. Currently, there is no cure for RP.

**[0004]** Gene therapy for RP was tested in proof-of-concept animal models, and later used as clinical treatment, where it improved vision in up-to half of the patients. In these trials, patients' genetic abnormalities were corrected by a gene supplementation approach (i.e., rescue via overexpression of a wild type (wt) gene). Initially, the gene supplementation appeared to work because patients experienced a functional rescue, but follow-up examination showed that degeneration of photoreceptors continued, and vision loss progressed in 3 years (Cideciyan et al. Proc Natl Acad Sci USA. 2013 Feb 5;110(6):E517-25; Bainbridge et al. N Engl J Med. 2015 May 14;372(20):1887-97; Jacobson et al. N Engl J Med. 2015 May 14;372(20):1920-6). Current gene therapy trials for other RP genes are also taking a gene supplementation approach and are likely to face similar hurdles unless the reasons for failure are addressed. Since supplementation with a wt gene leaves the patient's mutant gene intact, the presence of the mutant gene can continuously trigger ongoing damage despite the presence of a wt gene. The gene editing approach described in the present disclosure overcomes these and other obstacles to treatment of autosomal dominant as well as recessive diseases.

**[0005]** International patent application WO2014/204726 discloses the use of CRISPR-Cas9 systems to target genes in post-mitotic cells in specific tissues, e.g. retinal tissue. AAV-Spguides are disclosed.

## Summary of the Invention

**[0006]** The present invention provides a composition comprising at least one type of recombinant adeno-associated viral (AAV) vector encoding a CRISPR-Cas system directed to an autosomal dominant disease-related gene for use in treating an autosomal dominant ocular disease in a subject, the use comprising administering to the subject a therapeutically effective amount of the composition, wherein the at least one type of the AAV vector comprises:

1. (a) two guide RNA sequences that hybridize to the endogenous autosomal dominant disease-related gene in the subject;
2. (b) a second sequence comprising a codon-modified autosomal dominant disease-related gene or fragment thereof, wherein at least one disease related mutation has been corrected in the codon-modified autosomal dominant disease-related gene or fragment thereof, and wherein the codon-modified autosomal dominant disease-related gene or fragment thereof is not recognized by the guide RNA; and
3. (c) a third sequence encoding a Cas nuclease, optionally Cas 9,

wherein at least one guide RNA sequence is selected from the group consisting of SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7, or combinations thereof.

**[0007]** The endogenous autosomal dominant disease-related gene targeted by the present method may be wildtype and/or mutant.

**[0008]** A full-length or a fragment of a codon-modified autosomal dominant disease-related gene may be introduced into the subject according to the present invention.

**[0009]** In one embodiment, the two types of AAV vectors can be administered to the subject, where the first type of recombinant AAV vector comprises (i) the first sequence encoding at least one guide RNA and (ii) the second sequence comprising a codon-modified autosomal dominant disease-related gene or fragment thereof, and the second type of recombinant AAV vector comprises the third sequence, which encodes the Cas nuclease.

**[0010]** The ocular disease can include, but is not limited to, autosomal dominant chorioretinal atrophy or degeneration, autosomal dominant cone or cone-rod dystrophy, autosomal dominant congenital stationary night blindness, autosomal dominant leber congenital amaurosis, autosomal dominant macular degeneration, autosomal dominant ocular-retinal developmental disease, autosomal dominant optic atrophy, autosomal dominant retinitis pigmentosa, autosomal dominant syndromic/systemic diseases with retinopathy, sorsby macular dystrophy, age-related macular degeneration, doyne honeycomb macular disease, and juvenile macular degeneration.

**[0011]** In one embodiment, the ocular disease is retinitis pigmentosa. Retinitis pigmentosa can be caused by a mutation in *RHO* gene. The autosomal dominant disease-related gene may be the *RHO* gene. In another embodiment, the ocular disease is age-related macular degeneration. In a third embodiment, the ocular disease is doyne honeycomb. Doyme honeycomb may be caused by a mutation in the *EFEMP1* gene. The autosomal dominant disease-related gene may be the *EFEMP1*

gene.

**[0012]** The recombinant AAV vector may be an AAV2 vector. Alternatively, the AAV vector is an AAV8 vector. Other suitable AAV vectors may also be used.

**[0013]** The Cas nuclease can be Cas9. The CRISPR-Cas system can be under the control of a promoter which controls the expression of the codon-modified autosomal dominant disease-related gene product in ocular cells.

**[0014]** The codon-modified autosomal dominant disease-related gene sequence or fragment thereof may be integrated into the endogenous autosomal disease-related gene. Alternatively, the codon-modified autosomal dominant disease-related gene sequence or fragment is not integrated into the endogenous autosomal disease-related gene, but is present episomally.

**[0015]** The autosomal dominant disease-related gene may include, but is not limited to, PRDM13, RGR, TEAD1, AIPL1, CRX, GUCA1A, GUCY2D, PITPNM3, PROM1, PRPH2, RIMS1, SEMA4A, UNC119, GNAT1, PDE6B, RHO, WSF1, IMPDH1, OTX2, BEST1, C1QTNF5, CTNNA1, EFEMP1, ELOVL4, FSCN2, GUCA1B, HMCN1, IMPG1, RP1L1, TIMP 3, VCAN, MFN2, NR2F1, OPA1, ARL3, CA4, HK1, KLHL7, NR2E3, NRL, PRPF3, PRPF4, PRPF6, PRPF8, PRPF31, RDH12, ROM1, RP1, RP9, RPE65, SNRNP200, SPP2, TOPORS, ABCC6, ATXN7, COL11A1, COL2A1, JAG1, KCNJ13, KIF11, OPA3, PAX2, TREX1, CAPN5, CRB1, FZD4, ITM2B, LRP5, MAPKAPK3, MIR204, OPN1SW, RB1, TSPAN12, and ZNF408.

**[0016]** The recombinant AAV vector(s) may be administered by injection into the eye.

**[0017]** The codon-modified autosomal dominant disease-related gene or fragment thereof can be selected from the group consisting of, SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 23, SEQ ID NO: 24, SEQ ID NO: 25, SEQ ID NO: 26, SEQ ID NO: 27, or combinations thereof.

## BRIEF DESCRIPTION OF THE DRAWINGS

### **[0018]**

Figure 1A is schematic representations of the ChopStick AAV vectors. The left side shows a schematic representation of the AAV/Cas9 vector. Cas9 from *S. pyogenes* is driven by a 173-bp short CMV promoter (sCMV, SEQ ID NO: 14) and is terminated by a 50-bp synthetic poly-A signal (SPA) (SEQ ID NO: 19). The right side shows a schematic representation of the *RHO* sgRNAs and codon-modified cDNA (cm*RHO*, (SEQ ID NO: 9) expression vector. sgRNA1 and sgRNA2 are driven by U6 promoter (SEQ ID NO: 12). cm*RHO* cDNA with c-terminal tagged c-Myc is driven by CBh promoter (SEQ ID NO: 10) and terminated by bGH poly-A signal (SEQ ID NO: 11) Arrows indicate the direction of transcription. 5'- and 3'-ITR, inverted terminal repeats of AAV.

Figure 1B is a schematic representation of the ChopStick AAV gene therapy strategy. The left side schematic representation (I) shows that following co-infection of AAV/Cas9 and AAV/sgRNA1&2\_cm*RHO*, the co-expression of Cas9 protein and two h*RHO* exon 1-specific sgRNAs, sgRNA1 and sgRNA2, will lead to a 121- bp deletion in the host *RHO* Exon1. The right side

of the figure (II) shows the original and codon-modified rhodopsin sequence.

Figures 2A-B show that dual sgRNA provides more efficient "Chop" of *RHO* than single sgRNA. Figure 2A is a schematic representation of the target sites of sgRNA1 and sgRNA2 on *RHO*. The two sgRNAs both target *RHO* exon1, which is the beginning of the translation. Once the gene editing occurs, independent of whether one or two sgRNAs sites are targeted, most of the coding region will be affected. This design ensures the greatest disruption of gene expression and can be applied to many different types of *RHO* mutations. Figure 2B shows improved efficiency in truncating genes by the "Chop" strategy in human kidney cell line compared to using only one sgRNA. HEK293FT cells were transfected with Cas9 vector (pX459) carrying either no sgRNA, single sgRNA1, single sgRNA2, or both. Ninety-six hours later, DNA was extracted, and the *RHO* locus was amplified and analyzed by mismatch detection SURVEYOR assay. Applying two sgRNAs together resulted in gene deletion of approximately 30-40%, which indicated that "Chop" (gene deletion/disruption) strategy works efficiently in mammalian cells (lane 4). Using one sgRNA (lanes 2 and 3) at a time in contrast does not result in change in size of the *RHO* gene. Approximately 30% of the genomic DNA underwent non-homologous end joining (NHEJ) by one sgRNA. In contrast, up to 80% was edited (deletion and NHEJ) when two sgRNAs were used. AS a control, equal amounts of plasmid DNA (1  $\mu$ g /  $1 \times 10^5$  293FT cells) were used in each group.

Figures 3A-C show improved efficacy of inactivating a gene by dual sgRNA ("Chop" or gene deletion/disruption) when compared with a single sgRNA. Figure 3A is schematic representation of the target sites of sgRNA1 and sgRNA2 on a *RHO* expression vector. The two sgRNAs target the 5' end of *RHO* cDNA as indicated. Wt *RHO* cDNA was driven by a CMV promoter. EGFP driven by CMV promoter was used as an internal control in immunoblot assay, which normalizes the difference in transfection efficiency and protein loading. Figure 3B shows protein levels as measured by immunoblot when the HEK293FT cells were co-transfected with *RHO* expression vector and another vector expressing Cas9 machinery (pX459) carrying either sgRNA1, sgRNA2, or both. The sg3 group is a non-specific control sgRNA. Figure 3C indicates that, after normalization with EGFP, two sgRNAs together lower *RHO* expression by 70%, while using single sgRNA reduced expression only by 0-30% (compared to the control group (sg3)). This result indicated that "Chop" strategy can be used to significantly reduce or inactivate protein expression.

Figures 4A-C show that "Chop" (gene deletion or disruption strategy) has a potential to create a double strand break in order to facilitate precise repair through mechanism like homologous recombination. Figure 4A is schematic representation of the AAV-mediated CRISPR editing in *Rho*<sup>D190N</sup> mouse RP model. Dual virus treatment of AAV/Cas9 vector and a bicistronic AAV vector containing wt donor template and an sgRNA targeting D190N mutation would result in mutation-specific repair. Donor template construct contains wild-type *Rho* sequence with two modifications: 1) creation of an additional *Afl*III site upstream of the D190 codon for the identification of DNA replacement following CRISPR-induced homologous recombination and 2) introduction of 5 wobble base pairs (bps) to render the donor template unrecognizable by sgRNA and thus, Cas9-resistant. Figure 4B shows editing efficiency evaluated using tracking of indels (insertions and deletions) by decomposition (TIDE) analysis (publically available at <http://tide-calculator.nki.nl>: retrieved April 30, 2016) in mouse retina DNA treated with aforementioned AAV viruses, which showed that ~50% of photoreceptors underwent NHEJ. Figure 4C is a representative *Afl*III digestion of retinal DNA from a *Rho*<sup>D190N/+</sup> mouse showing a large portion of photoreceptors being repaired through homologous

recombination (lane 2). *Rho*<sup>D190N/+</sup> mice were treated with the Cas9 vector with (lane 2) or without (lane 1) the wild-type donor template, and retinal DNA was extracted and amplified with indicated screening primers.

Figures 5A-B show the histological and functional rescue by CRISPR/donor template-mediated repair. *Rho*<sup>D190N/+</sup> heterozygote mice were treated with dual virus treatment described in Figure 4A-C by subretinal injection at postnatal day 3. Figure 5B shows a visual function of mice evaluated by ERG following the treatment. Figure 5A shows a histological evaluation of the retinal tissue section. The H&E staining of retinal section shows the increase of photoreceptors (outernuclear layer, ONL) survival at 137%, as compared to the untreated eye (Figure 5A). The rectangular bars show an enlarged cross-sectional area of an ONL of photoreceptors in CRISPR/Cas9 (injected) and control eyes (untreated). The electroretinograms (ERGs) indicate a noticeable improvement in both a wave and b wave, of gene specific CRISPR-mediated therapy of 3-month old *Rho*<sup>D190N/+</sup> heterozygote (Figure 5B).

Figures 6A-C describe the generation of *RHO*-humanized animal model by CRISPR-mediated exon 1 replacement at the mouse *Rho* locus. This system enables the researchers to test CRISPR components *in vivo*. Figure 6A is an illustration of the strategy of replacing mouse (m) *Rho* exon 1 with either wild-type (wt) or mutant human (h) *RHO* exon 1. By co-electroporation of plasmid pX459 encoding Cas9 and *Rho* exon 1-specific sgRNA, a double strand break can be created in mouse exon1 that facilitates the homologous recombination in ES cells. Figure 6B shows restriction fragment length polymorphism (RFLP) assay results of ES cell DNA featuring additional *Avall* site indicating the replacement of mouse *Rho* exon 1 with human *RHO* exon 1 (lane 1 and 2). Figure 6C: Sequence electropherogram of PCR amplicons reveals fusion of human and mouse sequence from one targeted ES clone.

Figures 7A-C show the successful gene replacement of the D670G allele in the gene encoding *Pde6a* by CRISPR in mouse embryonic stem cells. Figure 7A is a schematic of donor construct which contains *Pde6a* with two changes: (1) a *Pde6a*-codon modification was introduced which creates an additional *SphI* site upstream from the D670G codon; and (2) eight wobble base pairs were introduced, making the donor template resistant to sgRNA targeting. Figure 7B shows PCR amplicons generated from ES cells that underwent homologous recombination. Figure 7C shows sequencing electropherogram data of target ES clone DNA, featuring an expected replacement of the D670G allele with donor template.

Figures 8A-C show that "ChopStick" (gene deletion or gene disruption) can be used to efficiently delete and correct a gene region of interest, such as one containing a mutation, in induced pluripotent stem (iPS) cell from a patient with juvenile macular degenerations (OMIM #126600). Figure 8A is a schematic illustration of the introduction of CRISPR components into human iPSCs. Cas9 protein/sgRNA complex (RNP) was co-nucleofected into human iPS cells with single strand donor template (ssODN). The clones were further selected and screened by restriction fragment length polymorphism (RFLP) assay. Figure 8B is a schematic representation of sgRNA targeting site in this case. The nucleotide marked with a dot corresponds to the mutation site. Figure 8C is the sequencing result of colony PCR, indicating replacement of donor template.

Figure 9 is a schematic representation of the self-excisional AAV/Cas9 vector. Cas9 from *S. pyogenes*, which is driven by a 173-bp short CMV promoter (sCMV) and terminated by a 50-bp

synthetic poly-A signal (SPA), is flanked by sgRNA-Y1 (SEQ ID NO: 7) target sequences (GGTTTTGGACAATGGAACCGTGG, originated from *Drosophila*). Once the cell expresses Cas9 protein and sgRNA-Y1 simultaneously, this AAV/Cas9 vector is destroyed by Cas9 itself.

## DETAILED DESCRIPTION

**[0019]** The term "nuclease" is used to generally refer to any enzyme that hydrolyzes nucleic acid sequences.

**[0020]** The term "ocular cells" refers to any cell in, or associated with the function of, the eye. The term may refer to any one or more of photoreceptor cells, including rod, cone and photosensitive ganglion cells, retinal pigment epithelium (RPE) cells, Müller cells, bipolar cells, horizontal cells, or amacrine cells. In one embodiment, the ocular cells are bipolar cells. In another embodiment, the ocular cells are horizontal cells. In yet a third embodiment, the ocular cells include ganglion cells.

**[0021]** The terms "polynucleotide", "nucleotide", "nucleotide sequence", "nucleic acid" and "oligonucleotide" are used interchangeably. These terms refer to a polymeric form of nucleotides of any length, either deoxyribonucleotides or ribonucleotides, or analogs. Examples of polynucleotides include, but are not limited to, coding or non-coding regions of a gene or gene fragment, exons, introns, messenger RNA (mRNA), transfer RNA, ribosomal RNA, short interfering RNA (siRNA), short-hairpin RNA (shRNA), micro-RNA (miRNA), ribozymes, cDNA, recombinant polynucleotides, branched polynucleotides, plasmids, vectors, isolated DNA of any sequence, isolated RNA of any sequence, nucleic acid probes, and primers. One or more nucleotides within a polynucleotide sequence can further be modified. The sequence of nucleotides may be interrupted by non-nucleotide components. A polynucleotide may also be modified after polymerization, such as by conjugation with a labeling agent.

**[0022]** The present disclosure is based, in part, on findings that gene editing can be used to correct the disease-causing mutant alleles, which in turn can be used for *in vivo* gene therapy for patients afflicted with autosomal dominant diseases.

**[0023]** The present disclosure takes advantage of the CRISPR gene-editing system, where the approach is to use a gene-editing enzyme with one or multiple unique single guide (sg) RNA sequences that target mutant allele(s) specifically or that target both the mutant and wild type alleles of a gene carrying an autosomal dominant mutation for destruction. This targeting is then followed by supplying the wild type gene cDNA, that is codon modified in order to evade recognition, by the sgRNA(s). Deletion of both the mutant and/or wild-type forms of the gene, followed by supplying the wild type gene cDNA that is codon modified and resistant to recognition by the guide RNAs results in the correction of the mutation, and thus, restoration of a phenotype found in the autosomal dominant diseases.

**[0024]** The inventors of the present disclosure refer to the gene-editing system described here as a "ChopStick" system. The "Chop" step involves partial or complete disruption of the i) mutant copy of

a gene that is to be corrected; and/or ii) the wild-type copy of said gene in a patient afflicted with autosomal dominant disease. Thus, the "Chop" step results in partial or complete loss of mutant and/or wild-type activity of the said gene. The "Stick" step encompasses the introduction of a codon-modified cDNA of a gene of interest or fragment thereof (characterized by the autosomal dominant mutation), which is intended to restore, correct, supplement, or augment the gene or gene product function in the cells.

**[0025]** In one embodiment, the "Stick" step results in integration of a codon-modified donor template of a gene of interest or fragment (characterized by the autosomal dominant mutation) into the endogenous autosomal disease-related gene. Such targeted integration is accomplished by homologous recombination. In general, a Cas-family nuclease makes a DNA double-strand break at a defined site in the genome, which can then be repaired by homologous recombination or non-homologous end joining.

**[0026]** Alternatively, the "Stick" step does not result in integration of a codon-modified donor template of a gene of interest or fragment (characterized by the autosomal dominant mutation) into the endogenous autosomal disease-related gene. For example, extrachromosomal, or episomal (episomally), vectors persist in the nucleus in an extrachromosomal state, and offer transgene delivery and expression without integration into the host genome. Among such vectors are AAV vectors, which are particularly efficient in transduction of nondividing cells, and where the vector genome persists predominantly in an episomal form.

**[0027]** A codon-modified donor template can be delivered to cells or a patient via episomal vectors. Because episomal vectors persist in multiple copies per cell, the resulting expression of the gene of interest may be comparatively high at both the RNA as well as protein level. In non-dividing cells, the presence of the AAV vector as an episomal replicating element may be sufficient for stable expression of the gene, RNA, and/or protein.

**[0028]** Given the general principles of the "Chop-Stick" system outlined in the present disclosure, the "Chop-Stick" system can be used as a gene-editing tool for the correction of the mutation(s) found in any autosomal dominant disease. Thus, the methods of the present disclosure can be used to treat any autosomal dominant disease, including, but not limited to, Acropectoral syndrome, Acute intermittent porphyria, Adermatoglyphia, Albright's hereditary osteodystrophy, Arakawa's syndrome II, Aromatase excess syndrome, Autosomal dominant cerebellar ataxia, Autosomal dominant retinitis pigmentosa, Axenfeld syndrome, Bethlem myopathy, Birt-Hogg-Dubé syndrome, Boomerang dysplasia, Branchio-oto-renal syndrome, Buschke-Ollendorff syndrome, Camurati-Engelmann disease, Central core disease, Collagen disease, Collagenopathy, types II and XI, Congenital distal spinal muscular atrophy, Congenital stromal corneal dystrophy, Costello syndrome, Currarino syndrome, Darier's disease, De Vivo disease, Dentatorubral-pallidoluysian atrophy, Dermatopathia pigmentosa reticularis, DiGeorge syndrome, Doyne honeycomb disease, Dysfibrinogenemia, Familial amyloid polyneuropathy, Familial atrial fibrillation, Familial hypercholesterolemia, Familial male-limited precocious puberty, Feingold syndrome, Felty's syndrome, Flynn-Aird syndrome, Gardner's syndrome, Gillespie syndrome, Gray platelet syndrome, Greig cephalopolysyndactyly syndrome, Hajdu-Cheney syndrome, Hawkinsinuria, Hay-Wells syndrome, Hereditary elliptocytosis, Hereditary hemorrhagic telangiectasia, Hereditary mucoepithelial dysplasia, Hereditary spherocytosis, Holt-Oram syndrome, Huntington's disease,

Hypertrophic cardiomyopathy, Hypoalphalipoproteinemia, Hypochondroplasia, Jackson-Weiss syndrome, Keratolytic winter erythema, Kniest dysplasia, Kostmann syndrome, Langer-Giedion syndrome, Larsen syndrome, Liddle's syndrome, Marfan syndrome, Marshall syndrome, Medullary cystic kidney disease, Metachondromatosis, Miller-Dieker syndrome, MOMO syndrome, Monilethrix, Multiple endocrine neoplasia, Multiple endocrine neoplasia type 1, Multiple endocrine neoplasia type 2, Multiple endocrine neoplasia type 2b, Myelokathexis, Myotonic dystrophy, Naegeli-Franceschetti-Jadassohn syndrome, Nail-patella syndrome, Noonan syndrome, Oculopharyngeal muscular dystrophy, Pachyonychia congenital, Pallister-Hall syndrome, PAPA syndrome, Papillorenal syndrome, Parastremmatic dwarfism, Pelger-Huet anomaly, Peutz-Jeghers syndrome, Polydactyly, Popliteal pterygium syndrome, Porphyria cutanea tarda, Pseudoachondroplasia, RASopathy, Reis-Bucklers corneal dystrophy, Romano-Ward syndrome, Rosselli-Gulienetti syndrome, Roussy-Lévy syndrome, Rubinstein-Taybi syndrome, Saethre-Chotzen syndrome, Schmitt Gillenwater Kelly syndrome, Short QT syndrome, Singleton Merten syndrome, Spinal muscular atrophy with lower extremity predominance, Spinocerebellar ataxia, Spinocerebellar ataxia type 6, Spondyloepiphyseal dysplasia congenital, Spondyloperipheral dysplasia, Stickler syndrome, Tietz syndrome, Timothy syndrome, Treacher Collins syndrome, Tuberous sclerosis, Upington disease, Variegate porphyria, Vitelliform macular dystrophy, Von Hippel-Lindau disease, Von Willebrand disease, Wallis-Zieff-Goldblatt syndrome, WHIM syndrome, White sponge nevus, Worth syndrome, Zaspopathy, Zimmermann-Laband syndrome, and Zori-Stalker-Williams syndrome. For example, in the Examples provided in the present disclosure, the inventors present the data performing "Chop" on human kidney cells and iPS cells (Figure 2 and Figure 8). These findings confirm the potential of the methods of the present disclosure to be used to prevent, correct, or treat autosomal dominant kidney diseases such as renal angiomyolipomas, medullary cystic kidney disease, or autosomal dominant polycystic kidney disease.

**[0029]** In further embodiments, the methods of the present disclosure can be used to prevent, correct, or treat ocular diseases that arise due to the presence of autosomal dominant mutation. Examples of such diseases include, but are not limited, autosomal dominant chorioretinal atrophy or degeneration, autosomal dominant cone or cone-rod dystrophy, autosomal dominant congenital stationary night blindness, autosomal dominant leber congenital amaurosis, autosomal dominant macular degeneration, autosomal dominant ocular-retinal developmental disease, autosomal dominant optic atrophy, autosomal dominant retinitis pigmentosa, autosomal dominant syndromic/systemic diseases with retinopathy, sorsby macular dystrophy, age-related macular degeneration, doyne honeycomb macular disease, and juvenile macular degeneration.

**[0030]** Generally, in the case of retinitis pigmentosa, patients with null rhodopsin mutations have a milder phenotype than those with severe dominant rhodopsin mutations. Even if the normal rhodopsin gene is destroyed together with the mutant one, the supply of the wild-type exon or cDNA (i.e., Stick) is still expected to improve retinal function in the recipient.

**[0031]** The methods of the present disclosure can be used for arresting progression of or ameliorating vision loss associated with retinitis pigmentosa (RP) in the subject. Vision loss linked to retinitis pigmentosa may include decrease in peripheral vision, central (reading) vision, night vision, day vision, loss of color perception, loss of contrast sensitivity, or reduction in visual acuity. The methods of the present disclosure can also be used to prevent, or arrest photoreceptor function loss, or increase photoreceptor function in the subject.

**[0032]** RP is diagnosed in part, through an examination of the retina. The eye exam usually reveals abnormal, dark pigment deposits that streak the retina. Additional tests for diagnosing RP include electroretinogram (ERG) and visual field testing.

**[0033]** Methods for measuring or assessing visual function, retinal function (such as responsiveness to light stimulation), or retinal structure in a subject are well known to one of skill in the art. See, e.g. Kanski's Clinical Ophthalmology: A Systematic Approach, Edition 8, Elsevier Health Sciences, 2015. Methods for measuring or assessing retinal response to light include may include detecting an electrical response of the retina to a light stimulus. This response can be detected by measuring an electroretinogram (ERG; for example full-field ERG, multifocal ERG, or ERG photostress test), visual evoked potential, or optokinetic nystagmus (see, e.g., Wester et al., Invest. Ophthalmol. Vis. Sci. 48:4542-4548, 2007). Furthermore, retinal response to light may be measured by directly detecting retinal response (for example by use of a microelectrode at the retinal surface). ERG has been extensively described by Vincent et al. Retina, 2013 Jan;33(1):5-12. Thus, methods of the present disclosure can be used to improve visual function, retinal function (such as responsiveness to light stimulation), retinal structure, or any other clinical symptoms or phenotypic changes associated with ocular diseases in subjects afflicted with ocular disease.

**[0034]** In one embodiment, the methods of the present disclosure can be used to prevent the development and progression of autosomal dominant disease. For example, a patient may be a carrier of autosomal dominant mutation, but the phenotypic expression of a disease has not been yet manifested, although the genomic defect has been identified by screening. The methods of the present disclosure may be applied to such patient to prevent the onset of disease.

**[0035]** Mutations in various genes have been identified to give rise to autosomal dominant diseases (such genes are also referred to as autosomal dominant disease-related genes). The methods of the present disclosure can be used to fully or partially correct mutations in such autosomal dominant disease-related genes, resulting in partial or full restoration of wild type.

**[0036]** In all cases where accession numbers are used, the accession numbers refer to one embodiment of the gene which may be used with the methods of the present disclosure. In one embodiment, the accession numbers are NCBI (National Center for Biotechnology Information) reference sequence (RefSeq) numbers.

**[0037]** For example, the autosomal dominant disease-related gene in retinitis pigmentosa may include, but are not limited to, ARL3(NC\_000010.11 (102673727..102714433, complement)), BEST1(NG\_009033.1), CA4(NG\_012050.1), CRX(NG\_008605.1), FSCN2(NG\_015964.1), GUCA1B(NG\_016216.1), HK1(NG\_012077.1), IMPDH1(NG\_009194.1), KLHL7(NG\_016983.1), NR2E3(NG\_009113.2), NRL(NG\_011697.1), PRPF3(NG\_008245.1), PRPF4(NG\_034225.1), PRPF6(NG\_029719.1), PRPF8(NG\_009118.1), PRPF31(NG\_009759.1), PRPH2(NG\_009176.1), RDH12(NG\_008321.1), RHO(NG\_009115.1), ROM1(NG\_009845.1), RP1(NG\_009840.1), RP9(NG\_012968.1), RPE65(NG\_008472.1), SEMA4A(NG\_027683.1), SNRNP200(NG\_016973.1), SPP2(NG\_008668.1), and TOPORS(NG\_017050.1). Genes and mutations causing autosomal dominant retinitis pigmentosa are in detail discussed by Daiger et al. (Cold Spring Harb Perspect Med. 2014 Oct 10;5(10)).

**[0038]** Another type of the autosomal dominant disease- related gene is autosomal dominant chorioretinal atrophy or degeneration-related gene, which may include:

PRDM13(NC\_000006.12 (99606774..99615578)), RGR(NG\_009106.1), and TEAD1(NG\_021302.1).

**[0039]** Another example of the autosomal dominant disease- related gene is autosomal dominant cone or cone-rod dystrophy-related gene, which can include: AIPL1(NG\_008474.1), CRX(NG\_008605.1), GUCA1A(NG\_009938.1), GUCY2D(NG\_009092.1), PITPNM3(NG\_016020.1), PROM1(NG\_011696.1), PRPH2(NG\_009176.1), RIMS1(NG\_016209.1), SEMA4A(NG\_027683.1), and UNC119(NG\_012302.1).

**[0040]** In one embodiment, the autosomal dominant disease- related gene is autosomal dominant congenital stationary night blindness-related gene, including:

GNAT1(NG\_009831.1), PDE6B(NG\_009839.1), and RHO(NG\_009115.1).

**[0041]** In another embodiment, the autosomal dominant disease- related gene is autosomal dominant deafness (alone or syndromic)-related gene such as WSF1(NC\_000004.12 (6269850..6303265)).

**[0042]** Another type of the autosomal dominant disease- related gene is autosomal dominant leber congenital amaurosis-related gene, which may include: CRX(NG\_008605.1), (NG\_009194.1), and OTX2(NG\_008204.1).

**[0043]** Another example of the autosomal dominant disease- related gene is autosomal dominant macular degeneration-related gene, which can include: BEST1(NG\_009033.1), C1QTNF5(NG\_012235.1), CTNNA1(NC\_000005.10 (138753396..138935034)), EFEMP1(NG\_009098.1), ELOVL4(NG\_009108.1), FSCN2(NG\_015964.1), GUCA1B(NG\_016216.1), HMCN1(NG\_011841.1), IMPG1(NG\_041812.1), OTX2(NG\_008204.1), PRDM13(NC\_000006.12 (99606774..99615578)), PROM1(NG\_011696.1), PRPH2(NG\_009176.1), RP1L1(NG\_028035.1, and TIMP3(NG\_009117.1).

**[0044]** In one embodiment, the autosomal dominant disease- related gene is autosomal dominant ocular retinal developmental disease-related gene such as VCAN(NG\_012682.1). The accession numbers are provided as specific examples of each gene which may be used with the methods of the disclosure.

**[0045]** In another embodiment, the autosomal dominant disease-related gene is autosomal dominant optic atrophy-related gene, including: MFN2(NG\_007945.1), NR2F1(NG\_034119.1), and OPA1(NG\_011605.1).

**[0046]** In one embodiment, the autosomal dominant disease-related gene is autosomal dominant syndromic/systemic disease with retinopathy-related gene, including: ABCC6(NG\_007558.2), ATXN7(NG\_008227.1), COL11A1(NG\_008033.1), COL2A1(NG\_008072.1), JAG1(NG\_007496.1), KCNJ13(NG\_016742.1), KIF11(NG\_032580.1), MFN2(NG\_007945.1), OPA3(NG\_013332.1), PAX2(NG\_008680.2), TREX1(NG\_009820.1), and VCAN(NG\_012682.1).

**[0047]** Another example of the autosomal dominant disease-related gene is autosomal dominant retinopathy-related gene, including: BEST1(NG\_009033.1), CAPN5(NG\_033002.1), CRB1(NG\_008483.2), FZD4(NG\_011752.1), ITM2B(NG\_013069.1), LRP5(NG\_015835.1), M APKAPK3(NC\_000003.12(50611862..50649297)),MIR204(NR\_029621.1), OPN1SW(NG\_009 094.1), RB1(NG\_009009.1), TSPAN12(NG\_023203.1), and ZNF408(NC\_000011.10 (46700767..46705916)).

**[0048]** In addition to being used for the prevention, correctness, or treatment of autosomal dominant diseases, the methods of the present disclosure can be used to prevent, correct, or treat any autosomal recessive diseases. Thus, all the methods described here as applicable to autosomal dominant diseases and autosomal dominant genes or fragments can be adopted for use in the treatment of autosomal recessive diseases.

**[0049]** In further embodiments, the methods of the present disclosure can be used to prevent, correct, or treat ocular diseases that arise due to the presence of autosomal recessive mutation. Examples of such diseases include, but are not limited to, autosomal recessive congenital stationary night, autosomal recessive deafness alone or syndromic, autosomal recessive leber congenital amaurosis, autosomal recessive optic atrophy, autosomal recessive retinitis pigmentosa, autosomal recessive syndromic/systemic diseases with retinopathy, autosomal recessive usher syndrome, other autosomal recessive retinopathy, autosomal recessive cone or cone-rod dystrophy, autosomal recessive macular degeneration, and autosomal recessive bardet-biedl syndrome.

**[0050]** According to the methods described here, autosomal recessive disease- related gene is corrected and can in-part or fully restore the function of a wild-type gene.

**[0051]** One type of the autosomal recessive disease- related gene is congenital stationary night - related gene, including: CABP4(NG\_021211.1), GNAT1(NG\_009831.1), GNB3(NG\_009100.1), GPR179(NG\_032655.2), GRK1(NC\_000013.11(113667279..113671659 )), GRM6(NG\_008105.1), LRIT3(NG\_033249.1), RDH5(NG\_008606.1),SAG(NG\_009116.1), SLC24A1(NG\_031968.2), and TRPM1(NG\_016453.2).

**[0052]** Another type of the autosomal recessive disease- related gene is bardet-biedl syndrome-related gene, including:

ADIPOR1(NC\_000001.1(202940825..202958572,complement)), ARL6(NG\_008119.2), BBIP1( NG\_041778.1), BBS1(NG\_009093.1), BBS2(NG\_009312.1), BBS4(NG\_009416.2), BBS5(NG\_ 011567.1), BBS7(NG\_009111.1), BBS9(NG\_009306.1),BBS10(NG\_016357.1), BBS12(NG\_02 1203.1), C8orf37(NG\_032804.1), CEP290(NG\_008417.1), IFT172(NG\_034068.1), IFT27(NG\_ 034205.1), INPP5E(NG\_016126.1), KCNJ13(NG\_016742.1),LZTFL1(NG\_033917.1), MKKS( NG\_009109.1), MKS1(NG\_013032.1), NPHP1(NG\_008287.1), SDCCAG8(NG\_027811.1), TRI M32(NG\_011619.1), and TTC8(NG\_008126.1).

**[0053]** One example of the autosomal recessive disease-related gene is cone or cone-rod dystrophy- related gene, including, but not limited to, ABCA4(NG\_009073.1), ADAM9(NG\_016335.1), ATF6(NG\_029773.1), C21orf2(NG\_032952.1), C8orf37(NG\_032804. 1), CACNA2D4(NG\_012663.1), CDHR1(NG\_028034.1),CERKL(NG\_021178.1), CNGA3(NG

\_009097.1), CNGB3(NG\_016980.1), CNNM4(NG\_016608.1), GNAT2(NG\_009099.1), KCNV2(NG\_012181.1), PDE6C(NG\_016752.1), PDE6H(NG\_016859.1), POC1B(NG\_041783.1), RAB28(NG\_033891.1), RAX2(NG\_011565.1), RDH5(NG\_008606.1), RPGRIP1(NG\_008933.1), and TTLL5(NG\_016974.1).

**[0054]** Another example of the autosomal recessive disease-related gene is deafness (alone or syndromic)-related gene including: CDH23(NG\_008835.1), CIB2(NG\_033006.1), DFNB31(NG\_016700.1), MYO7A(NG\_009086.1), PCDH15(NG\_009191.2), PDZD7(NG\_028030.1), and USH1C(NG\_011883.1).

**[0055]** In one embodiment, the autosomal recessive disease-related gene is leber congenital amaurosis-related gene, including: AIPL1(NG\_008474.1), CABP4(NG\_021211.1), CEP290(NG\_008417.1), CLUAP1(NG\_000016.10(3500945..3539048)), CRB1(NG\_008483.2), CRX(NG\_008605.1), DTHD1(NG\_032962.1), GDF6(NG\_008981.1), GUCY2D(NG\_009092.1), IFT140(NG\_032783.1), IQCB1(NG\_015887.1), KCNJ13(NG\_016742.1), LCA5(NG\_016011.1), LRAT(NG\_009110.1), NMNAT1(NG\_032954.1), PRPH2(NG\_009176.1), RD3(NG\_013042.1), RDH12(NG\_008321.1), RPE65(NG\_008472.1), RPGRIP1(NG\_008933.1), SPATA7(NG\_021183.1), and TULP1(NG\_009077.1).

**[0056]** In another embodiment, the autosomal recessive disease-related gene is optic atrophy-related gene, including: RTN4IP1(NG\_000006.12 (106571028..106630500, complement)), SLC25A46(NG\_000005.10(110738136..110765161)), and TMEM126A(NG\_017157.1).

**[0057]** One example of the autosomal recessive disease-related gene is retinitis pigmentosa-related gene, including: ABCA4(NG\_009073.1), AGLL5(NG\_000002.12 (27051423..27070622)), ARL6(NG\_008119.2), ARL2BP(NG\_033905.1), BBS1(NG\_009093.1), BBS2(NG\_009312.1), BEST1(NG\_009033.1), C2orf71(NG\_021427.1), C8orf37(NG\_032804.1), CERKL(NG\_021178.1), CLRN1(NG\_009168.1), CNGA1(NG\_009193.1), CNGB1(NG\_016351.1), CRB1(NG\_008483.2), CYP4V2(NG\_007965.1), DHDDS(NG\_029786.1), DHX38(NG\_034207.1), EMC1(NG\_032948.1), EYS(NG\_023443.2), FAM161A(NG\_028125.1), GPR125(NG\_000004.12 (22387374..22516058, complement)), HGSNAT(NG\_009552.1), IDH3B(NG\_012149.1), IFT140(NG\_032783.1), IFT172(NG\_034068.1), IMPG2(NG\_028284.1), KIAA1549(NG\_032965.1), KIZ(NG\_033122.1), LRAT(NG\_009110.1), MAK(NG\_030040.1), MERTK(NG\_011607.1), MVK(NG\_007702.1), NEK2(NG\_029112.1), NEUROD1(NG\_011820.1), NR2E3(NG\_009113.2), NRL(NG\_011697.1), PDE6A(NG\_009102.1), PDE6B(NG\_009839.1), PDE6G(NG\_009834.1), POMGNT1(NG\_009205.2), PRCD(NG\_016702.1), PROM1(NG\_011696.1), RBP3(NG\_029718.1), RGR(NG\_009106.1), RHO(NG\_009115.1), RLBP1(NG\_008116.1), RP1(NG\_009840.1), RP1L1(NG\_028035.1), RPE65(NG\_008472.1), SAG(NG\_009116.1), SLC7A14(NG\_034121.1), SPATA7(NG\_021183.1), TTC8(NG\_008126.1), TULP1(NG\_009077.1), USH2A(NG\_009497.1), ZNF408(NG\_000011.10 (46700767..46705916)), and ZNF513(NG\_028219.1).

**[0058]** Another example of the autosomal recessive disease-related gene is syndromic/systemic disease with retinopathy-related gene, including: ABCC6(NG\_007558.2), ABHD12(NG\_028119.1), ACBD5(NG\_032960.2), ADAMTS18(NG\_031879.1), ADIPOR1(NG\_000001.11(202940825..202958572, complement)), AHI1(NG\_008643.1), ALMS1(NG\_011690.1), CC2D2A(NG\_013035.1), CEP164(NG\_033032.1), CEP290(NG\_008417.1),

CLN3(NG\_008654.2), COL9A1(NG\_011654.1), CSPP1(NG\_034100.1), ELOVL4(NG\_00 9108.1), EXOSC2(NC\_000009.12 (130693760..130704894)), FLVCR1(NG\_028131.1), FLVCR1(NG\_028131.1), GNPTG(NG\_016985.1), HARS(NG\_032158.1), HGSNAT(NG\_0095 52.1), HMX1(NG\_013062.2), IFT140(NG\_032783.1), INPP5E(NG\_016126.1), INVS(NG\_0083 16.1), IQCB1(NG\_015887.1), LAMA1(NG\_034251.1), LRP5(NG\_015835.1), MKS1(NG\_0130 32.1), MTTTP(NG\_011469.1), NPHP1(NG\_008287.1), NPHP3(NG\_008130.1), NPHP4(NG\_011 724.2), OPA3(NG\_013332.1), PANK2(NG\_008131.3), PCYT1A(NG\_042817.1), PEX1(NG\_00 8341.1), PEX2(NG\_008371.1), PEX7(NG\_008462.1), PHYH(NG\_012862.1), PLK4(NG\_04182 1.1), PNPLA6(NG\_013374.1), POC1B(NG\_041783.1), PRPS1(NG\_008407.1), RDH11(NG\_04 2282.1), RPGRIP1L(NG\_008991.2), SDCCAG8(NG\_027811.1), SLC25A46(NC\_000005.10(11 0738136..110765161)), TMEM237(NG\_032049.1), TRNT1(NG\_041800.1), TTPA(NG\_016123 .1), TUB(NG\_029912.1), TUBGCP4(NG\_042168.1), TUBGCP6(NG\_032160.1), WDPCP(NG\_028144.1), WDR19(NG\_031813.1), WFS1(NG\_011700.1), and ZNF423(NG\_032972.2).

**[0059]** One type of the autosomal recessive disease- related gene is usher syndrome- related gene, including: ABHD12(NG\_028119.1), CDH23(NG\_008835.1), CEP250(NC\_000020.11(35455139..35517531)), CIB2(NG\_033006.1), CLRN1(NG\_009168.1), DFNB31(NG\_016700.1), GPR98(NG\_007083.1), HARS(NG\_032158.1), MYO7A(NG\_009086. 1), PCDH15(NG\_009191.2), USH1C(NG\_011883.1), USH1G(NG\_007882.1), and USH2A(NG\_009497.1).

**[0060]** Another type of the autosomal recessive disease- related gene is retinopathy- related gene, including: BEST1(NG\_009033.1), C12orf65(NG\_027517.1), CDH3(NG\_009096.1), CNGA3(NG\_009097.1), CNGB3(NG\_016980.1), CNNM4(NG\_016608.1), CYP4V2(NG\_0079 65.1), LRP5(NG\_015835.1), MFRP(NG\_012235.1), MVK(NG\_007702.1), NBAS(NG\_032964.1), NR2E3(NG\_009113.2), OAT(NG\_008861.1), PLA2G5(NG\_032045.1), PROM1(NG\_011696.1), RBP4(NG\_009104.1), RGS9(NG\_013021.1), RGS9BP(NG\_016751.1), and RLBP1(NG\_008116.1).

**[0061]** Yet another type of the autosomal recessive disease- related gene is macular degeneration- related gene, including: ABCA4(NG\_009073.1), CFH(NG\_007259.1), DRAM2(NC\_000001.11(111117332..111140216, complement)), IMPG1(NG\_041812.1), and MFSD8(NG\_008657.1).

**[0062]** In addition to being used for the prevention, correctness, or treatment of autosomal dominant and recessive diseases, the methods of the present disclosure can be used to prevent, correct, or treat any X-linked diseases. Thus, all the methods described here as applicable to autosomal dominant diseases and autosomal dominant genes or fragments can be adopted for use in the treatment of X-linked diseases.

**[0063]** Furthermore, the methods of the present disclosure can be used to prevent, correct, or treat ocular diseases that arise due to the presence of X-linked mutation. Examples of such diseases include: X-linked cone or cone-rod dystrophy, X-linked congenital stationary night blindness, X-linked macular degeneration, X-linked retinitis pigmentosa, X-linked syndromic/systemic diseases with retinopathy, X-linked optic atrophy, and X-linked retinopathies. According to the methods described here, X-linked disease- related gene is corrected and can in part or fully restore the function of a wild-type gene.

**[0064]** One example of the X-linked disease-related gene is cone or cone-rod dystrophy-related gene, including: CACNA1F(NG\_009095.2) and RPGR(NG\_009553.1).

**[0065]** Another example of the X-linked disease-related gene is congenital stationary night blindness-related gene, including: CACNA1F(NG\_009095.2) and NYX(NG\_009112.1).

**[0066]** In one embodiment, the X-linked disease-related gene is macular degeneration-related gene, such as RPGR(NG\_009553.1).

**[0067]** In another embodiment, the X-linked disease-related gene is optic atrophy-related gene, such as TIMM8A(NG\_011734.1).

**[0068]** One type of the X-linked disease-related gene is retinitis pigmentosa-related gene, including: OFD1(NG\_008872.1), RP2(NG\_009107.1), and RPGR(NG\_009553.1).

**[0069]** Another type of the X-linked disease-related gene is syndromic/systemic disease with retinopathy-related gene, including: OFD1(NG\_008872.1) and TIMM8A(NG\_011734.1).

**[0070]** Yet another example of the X-linked disease-related gene is retinopathy-related gene., including: CACNA1F(NG\_009095.2), CHM(NG\_009874.2), DMD(NG\_012232.1), NDP(NG\_009832.1), OPN1LW(NG\_009105.2), OPN1MW(NG\_011606.1), PGK1(NG\_008862.1), and RS1(NG\_008659.3).

**[0071]** In another embodiment, the methods of the present disclosure can be used to prevent, correct, or treat diseases that arise due to the presence of mutation in mitochondrial DNA. Such diseases may include, retinopathy caused by the gene mutations in mitochondrial DNA. Examples of genes that may be characterized by the mutation in mitochondrial DNA that causes the development of retinopathy include: MT-ATP6(NC\_012920.1 (8527..9207)), MT-TH(NC\_012920.1 (12138..12206)), MT-TL1(NC\_012920.1 (3230..3304)), MT-TP(NC\_012920.1 (15956..16023, complement), and MT-TS2(NC\_012920.1 (12207..12265)).

**[0072]** Table 1 provides an exemplary list of diseases and disease-related genes (accompanied with corresponding accession numbers) that can be treated and/or corrected using methods of the present disclosure.

**Table 1: Disease Related Disorders and Genes**

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Disease Category	Mapped and Identified Genes
Bardet-Biedl syndrome, autosomal recessive	ADIPOR1(NG_000001.11 (202940825..202958572, complement)), ARL6(NG_008119.2), BBIP1(NG_041778.1), BBS1(NG_009093.1), BBS2(NG_009312.1), BBS4(NG_009416.2), BBS5(NG_011567.1), BBS7(NG_009111.1), BBS9(NG_009306.1), BBS10(NG_016357.1), BBS12(NG_021203.1), C8orf37(NG_032804.1), CEP290(NG_008417.1), IFT172(NG_034068.1), IFT27(NG_034205.1), INPP5E(NG_016126.1), KCNJ13(NG_016742.1), LZTFL1(NG_033917.1), MKKS(NG_009109.1), MKS1(NG_013032.1), NPHP1(NG_008287.1), SDCCAG8(NG_027811.1), TRIM32(NG_011619.1), TTC8(NG_008126.1)
Chorioretinal atrophy or degeneration, autosomal dominant	PRDM13(NG_000006.12 (99606774..99615578)), RGR(NG_009106.1), TEAD1(NG_021302.1)
Cone or cone-rod dystrophy, autosomal dominant	AIPL1(NG_008474.1), CRX(NG_008605.1), GUCA1A(NG_009938.1), GUCY2D(NG_009092.1), PITPNM3(NG_016020.1), PROM1(NG_011696.1), PRPH2(NG_009176.1), RIMS1(NG_016209.1), SEMA4A(NG_027683.1), UNC119(NG_012302.1)
Cone or cone-rod dystrophy, autosomal recessive	ABCA4(NG_009073.1), ADAM9(NG_016335.1), ATF6(NG_029773.1), C21orf2(NG_032952.1), C8orf37(NG_032804.1), CACNA2D4(NG_012663.1), CDHR1(NG_028034.1), CERKL(NG_021178.1), CNGA3(NG_009097.1), CNGB3(NG_016980.1), CNNM4(NG_016608.1), GNAT2(NG_009099.1), KCNV2(NG_012181.1), PDE6C(NG_016752.1), PDE6H(NG_016859.1), POC1B(NG_041783.1), RAB28(NG_033891.1), RAX2(NG_011565.1), RDH5(NG_008606.1), RPGRIP1(NG_008933.1), TTLL5(NG_016974.1)
Cone or cone-rod dystrophy, X-linked	CACNA1F(NG_009095.2), RPGR(NG_009553.1)
Congenital stationary night blindness, autosomal dominant	GNAT1(NG_009831.1), PDE6B(NG_009839.1), RHO(NG_009115.1)
Congenital stationary night blindness, autosomal recessive	CABP4(NG_021211.1), GNAT1(NG_009831.1), GNB3(NG_009100.1), GPR179(NG_032655.2), GRK1(NG_000013.11 (113667279..113671659)), GRM6(NG_008105.1), LRIT3(NG_033249.1), RDH5(NG_008606.1), SAG(NG_009116.1), SLC24A1(NG_031968.2), TRPM1(NG_016453.2)
Congenital stationary night blindness, X-linked	CACNA1F(NG_009095.2), NYX(NG_009112.1)

Disease Category	Mapped and Identified Genes
Deafness alone or syndromic, autosomal dominant	WSF1(NC_000004.12 (6269850..6303265))
Deafness alone or syndromic, autosomal recessive	CDH23(NG_008835.1), CIB2(NG_033006.1), DFNB31(NG_016700.1), MYO7A(NG_009086.1), PCDH15(NG_009191.2), PDZD7(NG_028030.1), USH1C(NG_011883.1)
Leber congenital amaurosis, autosomal dominant	CRX(NG_008605.1), IMPDH1(NG_009194.1), OTX2(NG_008204.1)
Leber congenital amaurosis, autosomal recessive	AIPL1(NG_008474.1), CABP4(NG_021211.1), CEP290(NG_008417.1), CLUAP1(NC_000016.10 (3500945..3539048)), CRB1(NG_008483.2), CRX(NG_008605.1), DTHD1(NG_032962.1), GDF6(NG_008981.1), GUCY2D(NG_009092.1), IFT140(NG_032783.1), IQCB1(NG_015887.1), KCNJ13(NG_016742.1), LCA5(NG_016011.1), LRAT(NG_009110.1), NMNAT1(NG_032954.1), PRPH2(NG_009176.1), RD3(NG_013042.1), RDH12(NG_008321.1), RPE65(NG_008472.1), RPGRIP1(NG_008933.1), SPATA7(NG_021183.1), TULP1(NG_009077.1)
Macular degeneration, autosomal dominant	BEST1(NG_009033.1), C1QTNF5(NG_012235.1), CTNNA1(NC_000005.10 (138753396..138935034)), EFEMP1(NG_009098.1), ELOVL4(NG_009108.1), FSCN2(NG_015964.1), GUCA1B(NG_016216.1), HMCN1(NG_011841.1), IMPG1(NG_041812.1), OTX2(NG_008204.1), PRDM13(NC_000006.12 (99606774..99615578)), PROM1(NG_011696.1), PRPH2(NG_009176.1), RP1L1(NG_028035.1), TIMP3(NG_009117.1)
Macular degeneration, autosomal recessive	ABCA4(NG_009073.1), CFH(NG_007259.1), DRAM2(NC_000001.11(111117332..111140216, complement)), IMPG1(NG_041812.1), MFSD8(NG_008657.1)
Macular degeneration, X-linked	RPGR(NG_009553.1)
Ocular-retinal developmental disease, autosomal dominant	VCAN(NG_012682.1)

Disease Category	Mapped and Identified Genes
Optic atrophy, autosomal dominant	MFN2(NG_007945.1), NR2F1(NG_034119.1), OPA1(NG_011605.1 )
Optic atrophy, autosomal recessive	RTN4IP1(NC_000006.12(106571028..106630500, complement)), SLC25A46(NC_000005.10 (110738136..110765161)), TMEM126A(NG_017157.1)
Optic atrophy, X-linked	TIMM8A(NG_011734.1)
Retinitis pigmentosa, autosomal dominant	ARL3(NC_000010.11 (102673727..102714433, complement)), BEST1(NG_009033.1), CA4(NG_012050.1), CRX(NG_008605.1), FSCN2(NG_015964.1), GUCA1B(NG_016216.1), HK1(NG_012077.1), IMPDH1(NG_009194.1), KLHL7(NG_016983.1), NR2E3(NG_009113.2), NRL(NG_011697.1), PRPF3(NG_008245.1), PRPF4(NG_034225.1), PRPF6(NG_029719.1), PRPF8(NG_009118.1), PRPF31(NG_009759.1), PRPH2(NG_009176.1), RDH12(NG_008321.1), RHO(NG_009115.1), ROM1(NG_009845.1), RP1(NG_009840.1), RP9(NG_012968.1), RPE65(NG_008472.1), SEMA4A(NG_027683.1),
	SNRNP200(NG_016973.1), SPP2(NG_008668.1), TOPORS(NG_017050.1)
Retinitis pigmentosa, autosomal recessive	ABCA4(NG_009073.1), AGLB5(NC_000002.12 (27051423..27070622)), ARL6(NG_008119.2), ARL2BP(NG_033905.1), BBS1(NG_009093.1), BBS2(NG_009312.1), BEST1(NG_009033.1), C2orf71(NG_021427.1), C8orf37(NG_032804.1), CERKL(NG_021178.1), CLRN1(NG_009168.1), CNGA1(NG_009193.1), CNGB1(NG_016351.1), CRB1(NG_008483.2), CYP4V2(NG_007965.1), DHDHDS(NG_029786.1), DHX38(NG_034207.1), EMC1(NG_032948.1), EYS(NG_023443.2), FAM161A(NG_028125.1), GPR125(NC_000004.12 (22387374..22516058, complement)), HGSNAT(NG_009552.1), IDH3B(NG_012149.1), IFT140(NG_032783.1), IFT172(NG_034068.1), IMPG2(NG_028284.1), KIAA1549(NG_032965.1), KIZ(NG_033122.1), LRAT(NG_009110.1), MAK(NG_030040.1), MERTK(NG_011607.1), MVK(NG_007702.1), NEK2(NG_029112.1), NEUROD1(NG_011820.1), NR2E3(NG_009113.2), NRL(NG_011697.1), PDE6A(NG_009102.1), PDE6B(NG_009839.1), PDE6G(NG_009834.1), POMGNT1(NG_009205.2), PRC1(NG_016702.1), PROM1(NG_011696.1), RBP3(NG_029718.1), RGR(NG_009106.1), RHO(NG_009115.1), RLBP1(NG_008116.1), RP1(NG_009840.1), RP1L1(NG_028035.1), RPE65(NG_008472.1), SAG(NG_009116.1), SLC7A14(NG_034121.1), SPATA7(NG_021183.1), TTC8(NG_008126.1), TULP1(NG_009077.1), USH2A(NG_009497.1), ZNF408(NC_000011.10 (46700767..46705916)), ZNF513(NG_028219.1)
Retinitis	

Disease Category	Mapped and Identified Genes
pigmentosa, X-linked	OFD1(NG_008872.1), RP2(NG_009107.1, RPGR(NG_009553.1)
Syndromic/systemic diseases with retinopathy, autosomal dominant	ABCC6(NG_007558.2), ATXN7(NG_008227.1), COL11A1(NG_008033.1), COL2A1(NG_008072.1), JAG1(NG_007496.1), KCNJ13(NG_016742.1), KIF11(NG_032580.1), MFN2(NG_007945.1), OPA3(NG_013332.1), PAX2(NG_008680.2), TREX1(NG_009820.1), VCAN(NG_012682.1)
Syndromic/systemic diseases with retinopathy, autosomal recessive	ABCC6(NG_007558.2), ABHD12(NG_028119.1), ACBD5(NG_032960.2), ADAMTS18(NG_031879.1), ADIPOR1(NC_000001.11 (202940825..202958572,
	complement)), AHI1(NG_008643.1), ALMS1(NG_011690.1), CC2D2A(NG_013035.1), CEP164(NG_033032.1), CEP290(NG_008417.1), CLN3(NG_008654.2), COL9A1(NG_011654.1), CSPP1(NG_034100.1), ELOVL4(NG_009108.1), EXOSC2(NC_000009.12 (130693760..130704894)), FLVCR1(NG_028131.1), GNPTG(NG_016985.1), HARS(NG_032158.1), HGSNAT(NG_009552.1), HMX1(NG_013062.2), IFT140(NG_032783.1),
	INPP5E(NG_016126.1), INVS(NG_008316.1), IQCB1(NG_015887.1), LAMA1(NG_034251.1), LRP5(NG_015835.1), MKS1(NG_013032.1), MTTF(NG_011469.1), NPHP1(NG_008287.1), NPHP3(NG_008130.1), NPHP4(NG_011724.2), OPA3(NG_013332.1), PANK2(NG_008131.3), PCYT1A(NG_042817.1), PEX1(NG_008341.1), PEX2(NG_008371.1), PEX7(NG_008462.1),
	PHYH(NG_012862.1), PLK4(NG_041821.1), PNPLA6(NG_013374.1), POC1B(NG_041783.1), PRPS1(NG_008407.1), RDH11(NG_042282.1), RPGRIP1L(NG_008991.2), SDCCAG8(NG_027811.1), SLC25A46(NC_000005.10
Syndromic/systemic diseases with retinopathy, X-linked	(110738136..110765161)), TMEM237(NG_032049.1), TRNT1(NG_041800.1), TTPA(NG_016123.1), TUB(NG_029912.1), TUBGCP4(NG_042168.1), TUBGCP6(NG_032160.1), WDPCP(NG_028144.1), WDR19(NG_031813.1), WFS1(NG_011700.1), ZNF423(NG_032972.2)
Usher syndrome, autosomal recessive	OFD1(NG_008872.1), TIMM8A(NG_011734.1)
Usher syndrome, autosomal recessive	ABHD12(NG_028119.1), CDH23(NG_008835.1), CEP250(NC_000020.11 (35455139..35517531)), CIB2(NG_033006.1), CLRN1(NG_009168.1), DFNB31(NG_016700.1), GPR98(NG_007083.1), HARS(NG_032158.1),
	MYO7A(NG_009086.1), PCDH15(NG_009191.2), USH1C(NG_011883.1), USH1G(NG_007882.1), USH2A(NG_009497.1)

Disease Category	Mapped and Identified Genes
Other retinopathy, autosomal dominant	BEST1(NG_009033.1), CAPN5(NG_033002.1), CRB1(NG_008483.2), FZD4(NG_011752.1), ITM2B(NG_013069.1), LRP5(NG_015835.1), MAPKAPK3(NC_000003.12 (50611862..50649297)), MIR204(NR_029621.1), OPN1SW(NG_009094.1), RB1(NG_009009.1), TSPAN12(NG_023203.1), ZNF408(NC_000011.10 (46700767..46705916))
Other retinopathy, autosomal recessive	BEST1(NG_009033.1), C12orf65(NG_027517.1), CDH3(NG_009096.1), CNGA3(NG_009097.1), CNGB3(NG_016980.1), CNNM4(NG_016608.1), CYP4V2(NG_007965.1),
	LRP5(NG_015835.1), MFRP(NG_012235.1), MVK(NG_007702.1), NBAS(NG_032964.1), NR2E3(NG_009113.2), OAT(NG_008861.1), PLA2G5(NG_032045.1), PROM1(NG_011696.1), RBP4(NG_009104.1),
	RGS9(NG_013021.1), RGS9BP(NG_016751.1), RLBP1(NG_008116.1)
Other retinopathy, mitochondrial	MT-ATP6(NC_012920.1 (8527..9207)), MT-TH(NC_012920.1 (12138..12206)), MT-TL1(NC_012920.1 (3230..3304)), MT-TP(NC_012920.1 (15956..16023, complement)), MT-TS2(NC_012920.1 (12207..12265))
Other retinopathy, X-linked	CACNA1F(NG_009095.2), CHM(NG_009874.2), DMD(NG_012232.1), NDP(NG_009832.1), OPN1LW(NG_009105.2), OPN1MW(NG_011606.1), PGK1(NG_008862.1), RS1(NG_008659.3)

**[0073]** The methods of the present disclosure can also be used to prevent, correct, or treat cancers that arise due to the presence of mutation in a tumor suppressor gene. Examples of tumor suppression genes include: retinoblastoma susceptibility gene (RB) gene, p53 gene, deleted in colon carcinoma (DCC) gene, adenomatous polyposis coli (APC) gene, p16, BRCA1, BRCA2, MSH2, and the neurofibromatosis type 1 (NF-1) tumor suppressor gene (Lee et al. Cold Spring Harb Perspect Biol. 2010 Oct; 2(10):).

**[0074]** Tumor suppressor genes are genes that, in their wild-type alleles, express proteins that suppress abnormal cellular proliferation. When the gene coding for a tumor suppressor protein is mutated or deleted, the resulting mutant protein or the complete lack of tumor suppressor protein expression may fail to correctly regulate cellular proliferation, and abnormal cellular proliferation may take place, particularly if there is already existing damage to the cellular regulatory mechanism. A number of well-studied human tumors and tumor cell lines have been shown to have missing or nonfunctional tumor suppressor genes. Thus, a loss of function or inactivation of tumor suppressor genes may play a central role in the initiation and/or progression of a significant number of human cancers.

**[0075]** The methods of the present disclosure may be used to treat patients at a different stage of the disease (e.g. early, middle or late). The present methods may be used to treat a patient once or

multiple times. Thus, the length of treatment may vary and may include multiple treatments.

**[0076]** As discussed in the present disclosure, the methods of the present disclosure can be used for correcting or treating autosomal dominant ocular disease in a subject. For example, the "Chop" step involves deletion of both the mutant copy of the autosomal dominant ocular disease-related gene that is to be corrected, and/or the endogenous wild-type copy of the same gene in a patient afflicted with autosomal dominant ocular disease. Thus, the "Chop" step results in complete or partial loss of both mutant and/or wild-type activity of a gene. The autosomal dominant ocular disease-related gene is then corrected using the "Stick" step, which involves the introduction of a sequence encoding a modified autosomal dominant ocular disease-related gene or fragment. The modified, autosomal dominant ocular disease-related gene sequence can be modified in such a way that it is not recognized (unrecognizable) by sgRNA, which targets the wild-type or mutant form of the gene (non-codon-modified form of the gene). This modification renders the codon-modified donor template resistant to the Cas-family nuclease.

**[0077]** The constructs encoding the "Chop" and "Stick" components can be delivered to the subject using one or more recombinant adeno-associated viral (AAV) vectors (e.g., 1, 2, 3, 4, 5, 6, 7, 8, 9, or more AAV vectors). One or more sgRNAs can be packaged into single (one) recombinant AAV vector. The recombinant AAV vector may also include codon-modified autosomal dominant ocular disease-related gene sequence (donor template). A Cas-family nuclease can be packaged into the same, or alternatively separate recombinant AAV vectors.

**[0078]** The method described here also provides for correcting autosomal dominant ocular disease in a subject, comprising administering to said subject by injection a therapeutically effective amount of a recombinant AAV virus encoding a nucleic acid sequence comprising a CRISPR system polynucleotide sequence, wherein the polynucleotide sequence comprises: (i) one or more guide RNA sequences that hybridize to an autosomal dominant disease-related gene sequence; (ii) a second sequence encoding a codon-modified autosomal dominant disease-related gene or fragment, wherein at least one disease related mutation in the modified autosomal dominant disease-related gene or fragment has been corrected and the codon-modified autosomal dominant disease related gene or fragment cannot be recognized by one or more sg RNA sequences that hybridize to an unmodified autosomal dominant disease-related gene sequence; and (iii) a sequence encoding a Cas family enzyme.

**[0079]** As the carrying capacity of AAV may pose challenges, two or more AAV vectors can be used simultaneously. For example, a Cas family nuclease may be packaged into a different AAV vectors. Furthermore, sequences encoding sgRNA(s), codon-modified autosomal dominant disease-related gene or fragment, and a Cas family nuclease can each be packaged into a separate AAV vector.

**[0080]** In the case of RP treatment, the methods of the present disclosure can comprise: administering to a subject by injection a therapeutically effective amount of a (1) recombinant AAV virus encoding a nucleic acid sequence comprising a CRISPR system polynucleotide sequence, wherein the polynucleotide sequence comprises: (i) two guide RNA sequences that hybridize to mutant and wild type RHO sequences; (ii) a second sequence encoding a codon-modified RHO gene or fragment, where the mutation(s) of the endogenous RHO gene has been corrected and the modified RHO gene or fragment cannot be recognized by one or more sgRNA sequences that

hybridize to the mutant and wild type RHO gene sequence; and (2) a second recombinant AAV virus encoding a Cas family enzyme.

**[0081]** Generally, co-expression of a Cas-family enzyme and an autosomal dominant disease-related gene-specific sgRNAs in ocular cells, leads to truncation of the autosomal dominant disease-related gene, thereby preventing the expression of either the wild-type (wt) or disease-causing mutant gene. Simultaneously, codon-modified cDNA of the autosomal dominant disease-related gene may also be supplied to ocular cells, where the coding sequence of autosomal dominant disease-related gene is modified in such a way that is resistant to sgRNAs (and thus resistant to Cas family nuclease). This strategy results in the expression of codon-modified cDNA of the autosomal dominant disease-related gene, which can restore or correct the function of the autosomal dominant disease-related gene or fragment after the deletion of endogenous gene(s) or fragments.

**[0082]** The codon-modified cDNA (donor-template) may be modified in such a way as to render it unrecognizable by the sgRNA(s) used to target either mutant and wt disease-related gene(s). Thus, mutations need to be introduced into a donor-template gene or fragment to avoid this donor-template gene or fragment being recognized by sgRNA(s) and consequently degraded by Cas enzyme (for example a Cas9 nuclease) which has been introduced in cells. This can be accomplished by introducing a wobble base into donor-template, thus making sure that the change in DNA results in a silent mutation, leaving the expression product of wt gene intact. The term "wobble base" as used in the present disclosure refers to a change in a one or more nucleotide bases of a reference nucleotide sequence wherein the change does not change the sequence of the amino acid coded by the nucleotide relative to the reference sequence.

**[0083]** The number of wobble bases that need to be introduced into donor-template may range from about 1-30, about 1-20, about 2-19, about 3-18, about 4-17, about 5-16, about 6-15, about 7-14, about 8-13, about 9-12, about 10-11, about 9, about 8, about 7, about 6., or about 5. Additionally, given the numerous software applications available for in-silico predictive modeling of sgRNA, one can perform in-silico analysis to test whether codon-modified donor-template would be recognized by sgRNA. An example of publically available CRISPR sgRNA tool can be found at <http://www.genscript.com/gRNA-design-tool.html>: retrieved April 30, 2016.

**[0084]** Alternatively, if the goal of the treatment is to delete, destroy, or truncate only mutated form of a gene or a fragment, and leave the wild type form intact, donor template or wild type gene sequence that is supplemented to the cells or a patient may not be codon-modified. Under such circumstances, sgRNA(s) used as part of the CRISPR components would be designed to recognize and target only the mutated form of a disease-related gene (and not recognize and target a wild type (such as donor-template) form of said gene).

**[0085]** The methods of the present disclosure have been applied to various genes, including PDE6A, EFEMP1, mouse Rhodopsin (RHO), and human RHO genes. RP can be caused by autosomal recessive mutations in the PDE6A gene, or autosomal dominant mutations in RHO gene. Mutations in EFEMP1 are responsible for autosomal dominant Malattia Leventinese (ML) and Doyme honeycomb retinal dystrophy (DHRD). Moreover, the methods have been applied to various cell types, including, but not limited to, mouse retina cells as well as human iPS cells. Additionally,

the methods described here have also been applied *in vivo* using a mouse model of ocular disease. Thus, methods of the present disclosure can be applied to both animal as well as human subjects.

**[0086]** Furthermore, methods of the present disclosure that have been applied to specific gene-humanized mouse model as well as patient-derived cells allow for determining the efficiency and efficacy of designed sgRNA and site-specific recombination frequency in human cells, which can be then used as a guide in a clinical setting.

**[0087]** In one embodiment, the "ChopStick" system comprises the following components: two recombinant AAV vectors: the first carrying a polynucleotide encoding the Cas9 enzyme to "Chop" the mutant and/or native rhodopsin genes, and the second carrying a nucleotide encoding the codon-modified human rhodopsin cDNA to "Stick" the normal rhodopsin back into the patient. The codon-modified or genetically engineered human rhodopsin sequence, which is driven by the CBh promoter is resistant to destruction by the gene-editing enzyme, rescues the patient's phenotype.

**[0088]** In one embodiment, the present method provides at least 50% rhodopsin levels from the CBh promoter-driven codon-modified RHO cDNA, which are sufficient to improve survival. In another embodiment, there is not an excessive amount of rhodopsin expressed using the codon-modified *RHO* donor sequence.

**[0089]** For studies using human and patient-derived cells, the inventors chose AAV2 vector as a backbone vector for all the constructs, as it has been shown that AAV2 may transduce human iPS more efficiently than other AAV vectors (Mitsui K et al. *Biochem Biophys Res Commun.* 2009 Oct 30;388(4):711-7; Deyle DR et al. *Mol Ther.* 2012 Jan;20(1):204-13; and Deyle DR et al. *Nucleic Acids Res.* 2014 Mar;42(5):3119-24). However, a variety of other AAV vectors may also be used to carry out the methods of the present disclosure.

**[0090]** The degree of improvement of the autosomal dominant disease by the present methods can vary. For example, the present methods may restore about 20%, about 30%, about 40%, greater than 10%, greater than 20%, greater than 30%, greater than 40%, greater than 50%, greater than 60%, greater than 70%, greater than 80%, or greater than 90%, of the autosomal dominant disease-related gene expression, of the normal levels of the gene product in a control subject, which may be age and sex matched.

**[0091]** In certain embodiments, expression of a wild-type gene (e.g., rhodopsin) can be observed in about 2 weeks following administration to a subject and/or cells. Expression may be maintained for unlimited period of time in nondividing somatic cells (e.g., photoreceptors, neuron cells, muscle cells, etc.). In one embodiment, expression of wild-type rhodopsin is observed in about 3 days, in about 1 week, in about 3 weeks, in about 1 month, in about 2 months, from about 1 week to about 2 weeks, or within different time-frames.

**[0092]** According to the various embodiments of the present disclosure, a variety of known viral constructs may be used to deliver desired (Chop and Stick) components such as Cas-family nuclease, sgRNA(s), codon-modified wild-type gene (also referred to as codon-modified donor template), donor template, etc. to the targeted cells and/or a subject. Nonlimiting examples of such recombinant viruses include recombinant adeno-associated virus (AAV), recombinant

adenoviruses, recombinant lentiviruses, recombinant retroviruses, recombinant poxviruses, and other known viruses in the art, as well as plasmids, cosmids, and phages. Options for gene delivery viral constructs are well known (see, e.g., Ausubel et al., *Current Protocols in Molecular Biology*, John Wiley & Sons, New York, 1989; Kay, M. A., et al., 2001 *Nat. Medic.* 7(1):33-40; and Walther W. and Stein U., 2000 *Drugs*, 60(2): 249-71).

**[0093]** Additionally delivery vehicles such as nanoparticle- and lipid-based mRNA or protein delivery systems can be used as an alternative to AAV vectors. Further examples of alternative delivery vehicles include lentiviral vectors, lipid-based delivery system, gene gun, hydrodynamic, electroporation or nucleofection microinjection, and biolistics. Various gene delivery methods are discussed in detail by Nayerossadat et al. (*Adv Biomed Res.* 2012; 1: 27) and Ibraheem et al. (*Int J Pharm.* 2014 Jan 1;459(1-2):70-83).

**[0094]** The present methods may utilize adeno-associated virus (AAV) mediated genome engineering. AAV vectors possess a broad host range; transduce both dividing and non-dividing cells *in vitro* and *in vivo* and maintain high levels of expression of the transduced genes. Viral particles are heat stable, resistant to solvents, detergents, changes in pH, temperature, and can be concentrated on CsCl gradients. AAV is not associated with any pathogenic event, and transduction with AAV vectors has not been found to induce any lasting negative effects on cell growth or differentiation. In contrast to other vectors, such as lentiviral vectors, AAVs lack integration machinery and have been approved for clinical use (Wirth et al. *Gene.* 2013 Aug 10;525(2): 162-9).

**[0095]** The single-stranded DNA AAV viral vectors have high transduction rates in many different types of cells and tissues. Upon entering the host cells, the AAV genome is converted into double-stranded DNA by host cell DNA polymerase complexes and exist as an episome. In non-dividing host cells, the episomal AAV genome can persist and maintain long-term expression of a therapeutic transgene. (*J Virol.* 2008 Aug; 82(16): 7875-7885).

**[0096]** AAV vectors and viral particles of the present disclosure may be employed in various methods and uses. In one embodiment, a method encompasses delivering or transferring a heterologous polynucleotide sequence into a patient or a cell of a patient and includes administering a viral AAV particle, a plurality of AAV viral particles, or a pharmaceutical composition of a AAV viral particle or plurality of AAV viral particles to a patient or a cell of the patient, thereby delivering or transferring a heterologous polynucleotide sequence into the patient or cell of the patient.

**[0097]** In another embodiment, the method is for treating a patient deficient or in need of protein expression or function, or in need of reduced expression or function of an endogenous protein (e.g., an undesirable, aberrant or dysfunctional protein), that includes providing a recombinant AAV viral particle, a plurality of recombinant AAV viral particles, or a pharmaceutical composition of a recombinant AAV viral particle or plurality of AAV viral particles; and administering the recombinant AAV viral particle, plurality of recombinant AAV viral particles, or pharmaceutical composition of AAV viral particle or plurality of AV viral particles to the patient, where the heterologous polynucleotide sequence is expressed in the patient, or wherein the heterologous polynucleotide sequence encodes one or more sgRNA(s) that reduces and or deletes endogenous DNA segment (e.g., an undesirable, aberrant or dysfunctional DNA segment) in the patient, and where the heterologous

polynucleotide sequence encodes a codon modified gene or fragment thereof that is not recognizable by one or more sgRNA(s) used to reduce and or delete endogenous DNA segment.

**[0098]** The characterization of new AAV serotypes has revealed that they have different patterns of transduction in diverse tissues. For illustrative purposes, AAV2 and AAV8 were used in the Examples of the present disclosure; however, for the purposes of the present invention, AAV viral vectors may be selected from among any AAV serotype, including, without limitation, AAV1, AAV2, AAV3, AAV4, AAV5, AAV6, AAV7, AAV8, AAV9, AAV10 or other known and unknown AAV serotypes.

**[0099]** The term AAV covers all subtypes, serotypes and pseudotypes, and both naturally occurring and recombinant forms, except where required otherwise. Pseudotyped AAV refers to an AAV that contains capsid proteins from one serotype and a viral genome of a second serotype.

**[0100]** To minimize the intensity and duration of Cas9 expression and potential off-targeting effects, self-excisional AAV-Cas9 vectors have also been generated, which have the ability to self-inactivate Cas9 expression shortly after Cas9 production. This approach comprises flanking the Cas9 gene with two sgRNA-Y1 target sites (similar to loxP sites in Cre recombinase system) to terminate Cas9 own expression (as shown in Figure 9). It is anticipated that the amount of Cas9 enzyme present (before it terminates itself) is still sufficient to cut the desired locus (such as Rho or PDE6A locus for example).

**[0101]** The design of self-inactivating recombinant AAV vectors (see Figure 9) enables the inventors to control the amount and duration of Cas9 expression in target cells, and can prevent the unwanted off-target effects due to excessive expression of Cas9 protein.

**[0102]** Vectors of the present disclosure can comprise any of a number of promoters known to the art, wherein the promoter is constitutive, regulatable or inducible, cell type specific, tissue-specific, or species specific. In addition to the sequence sufficient to direct transcription, a promoter sequence of the invention can also include sequences of other regulatory elements that are involved in modulating transcription (e.g., enhancers, kozak sequences and introns). Many promoter/regulatory sequences useful for driving constitutive expression of a gene are available in the art and include, but are not limited to, for example, CMV (cytomegalovirus promoter), EF1 $\alpha$  (human elongation factor 1 alpha promoter), SV40 (simian vacuolating virus 40 promoter), PGK (mammalian phosphoglycerate kinase promoter), Ubc (human ubiquitin C promoter), human beta-actin promoter, rodent beta-actin promoter, CB $\beta$  (chicken beta-actin promoter), CAG (hybrid promoter contains CMV enhancer, chicken beta actin promoter, and rabbit beta-globin splice acceptor), TRE (Tetracycline response element promoter), H1 (human polymerase III RNA promoter), U6 (human U6 small nuclear promoter), and the like. Moreover, inducible and tissue specific expression of an RNA, transmembrane proteins, or other proteins can be accomplished by placing the nucleic acid encoding such a molecule under the control of an inducible or tissue specific promoter/regulatory sequence. Examples of tissue specific or inducible promoter/regulatory sequences which are useful for this purpose include, but are not limited to, the rhodopsin promoter, the MMTV LTR inducible promoter, the SV40 late enhancer/promoter, synapsin 1 promoter, ET hepatocyte promoter, GS glutamine synthase promoter and many others. Various commercially available ubiquitous as well as tissue-specific promoters can be found at <http://www.invivogen.com/prom-a-list>. In addition, promoters which are well known in the art can be

induced in response to inducing agents such as metals, glucocorticoids, tetracycline, hormones, and the like, are also contemplated for use with the invention. Thus, it will be appreciated that the present disclosure includes the use of any promoter/regulatory sequence known in the art that is capable of driving expression of the desired protein operably linked thereto.

**[0103]** Vectors according to the present disclosure can be transformed, transfected or otherwise introduced into a wide variety of host cells. Transfection refers to the taking up of a vector by a host cell whether or not any coding sequences are in fact expressed. Numerous methods of transfection are known to the ordinarily skilled artisan, for example, lipofectamine, calcium phosphate coprecipitation, electroporation, DEAE-dextran treatment, microinjection, viral infection, and other methods known in the art. Transduction refers to entry of a virus into the cell and expression (e.g., transcription and/or translation) of sequences delivered by the viral vector genome. In the case of a recombinant vector, "transduction" generally refers to entry of the recombinant viral vector into the cell and expression of a nucleic acid of interest delivered by the vector genome.

**[0104]** The method of treating an autosomal dominant ocular disease in a patient can comprise administering to the patient an effective concentration of a composition comprising any of the recombinant AAVs described herein and a pharmaceutically acceptable carrier. In one embodiment, an effective concentration of virus is  $1 \times 10^6$  -  $11 \times 10^{13}$  GC/ml (genome copies/ml). The range of viral concentration effective for the treatment can vary depending on factors including, but not limited to specific mutation, patient's age, and other clinical parameters.

**[0105]** Recombinant AAV vectors(s) encoding CRISPR-Cas components and/or codon-modified donor-template comprising autosomal dominant disease-related gene or fragment can be produced *in vitro*, prior to administration into a patient. Production of recombinant AAV vectors and their use in *in vitro* and *in vivo* administration has been discussed in detail by Gray et al. (Curr. Protoc. Neurosci. 2011 Oct, Chapter:Unit 4.17).

**[0106]** The recombinant AAV containing the desired recombinant DNA can be formulated into a pharmaceutical composition intended for subretinal or intravitreal injection. Such formulation involves the use of a pharmaceutically and/or physiologically acceptable vehicle or carrier, particularly one suitable for administration to the eye, e.g., by subretinal injection, such as buffered saline or other buffers, e.g., HEPES, to maintain pH at appropriate physiological levels, and, optionally, other medicinal agents, pharmaceutical agents, stabilizing agents, buffers, carriers, adjuvants, diluents, etc. For injection, the carrier will typically be a liquid. Exemplary physiologically acceptable carriers include sterile, pyrogen-free water and sterile, pyrogen-free, phosphate buffered saline.

**[0107]** In one embodiment, the carrier is an isotonic sodium chloride solution. In another embodiment, the carrier is balanced salt solution. In one embodiment, the carrier includes tween. If the virus is to be stored long-term, it may be frozen in the presence of glycerol or Tween-20. In another embodiment, the pharmaceutically acceptable carrier comprises a surfactant, such as perfluorooctane (Perfluron liquid). In certain embodiments, the pharmaceutical composition described above is administered to the subject by subretinal injection. In other embodiments, the pharmaceutical composition is administered by intravitreal injection. Other forms of administration that may be useful in the methods described herein include, but are not limited to, direct delivery to

a desired organ (e.g., the eye), oral, inhalation, intranasal, intratracheal, intravenous, intramuscular, subcutaneous, intradermal, and other parental routes of administration. Additionally, routes of administration may be combined, if desired.

**[0108]** In preferred embodiments, route of administration is subretinal injection or intravitreal injection.

**[0109]** Methods for modification of genomic DNA are well known in the art. For example, methods may use a DNA digesting agent to modify the DNA by either the non-homologous end joining DNA repair pathway (NHEJ) or the homology directed repair (HDR) pathway. The term "DNA digesting agent" refers to an agent that is capable of cleaving bonds (i.e. phosphodiester bonds) between the nucleotide subunits of nucleic acids.

**[0110]** In one embodiment, the DNA digesting agent is a nuclease. Nucleases are enzymes that hydrolyze nucleic acids. Nucleases may be classified as endonucleases or exonucleases. An endonuclease is any of a group of enzymes that catalyze the hydrolysis of bonds between nucleic acids in the interior of a DNA or RNA molecule. An exonuclease is any of a group of enzymes that catalyze the hydrolysis of single nucleotides from the end of a DNA or RNA chain. Nucleases may also be classified based on whether they specifically digest DNA or RNA. A nuclease that specifically catalyzes the hydrolysis of DNA may be referred to as a deoxyribonuclease or DNase, whereas a nuclease that specifically catalyzes the hydrolysis of RNA may be referred to as a ribonuclease or an RNase. Some nucleases are specific to either single-stranded or double-stranded nucleic acid sequences. Some enzymes have both exonuclease and endonuclease properties. In addition, some enzymes are able to digest both DNA and RNA sequences.

**[0111]** Non-limiting examples of the endonucleases include a zinc finger nuclease (ZFN), a ZFN dimer, a ZFNickase, a transcription activator-like effector nuclease (TALEN), or a RNA-guided DNA endonuclease (e.g., CRISPR/Cas9). Meganucleases are endonucleases characterized by their capacity to recognize and cut large DNA sequences (12 base pairs or greater). Any suitable meganuclease may be used in the present methods to create double-strand breaks in the host genome, including endonucleases in the LAGLIDADG and PI-Sce family.

**[0112]** One example of a sequence-specific nuclease system that can be used with the methods and compositions described herein includes the CRISPR system (Wiedenheft, B. et al. *Nature* 482, 331-338 (2012); Jinek, M. et al. *Science* 337, 816-821 (2012); Mali, P. et al. *Science* 339, 823-826 (2013); Cong, L. et al. *Science* 339, 819-823 (2013)). The CRISPR (Clustered Regularly Interspaced Short Palindromic Repeats) system exploits RNA-guided DNA-binding and sequence-specific cleavage of target DNA. The guide RNA/Cas combination confers site specificity to the nuclease. A single guide RNA (sgRNA) contains about 20 nucleotides that are complementary to a target genomic DNA sequence upstream of a genomic PAM (protospacer adjacent motifs) site (NGG) and a constant RNA scaffold region. The Cas (CRISPR-associated) protein binds to the sgRNA and the target DNA to which the sgRNA binds and introduces a double-strand break in a defined location upstream of the PAM site. Cas9 harbors two independent nuclease domains homologous to HNH and RuvC endonucleases, and by mutating either of the two domains, the Cas9 protein can be converted to a nickase that introduces single-strand breaks (Cong, L. et al. *Science* 339, 819-823 (2013)). It is specifically contemplated that the methods and compositions of

the present disclosure can be used with the single- or double-strand-inducing version of Cas9, as well as with other RNA-guided DNA nucleases, such as other bacterial Cas9-like systems. The sequence-specific nuclease of the present methods and compositions described herein can be engineered, chimeric, or isolated from an organism. The nuclease can be introduced into the cell in form of a DNA, mRNA and protein.

**[0113]** In one embodiment, the methods of the present disclosure comprise using one or more sgRNAs to "Chop", remove, or suppress an autosomal dominant disease- related gene. In another embodiment, one sgRNA(s) is used to "Chop", remove, or suppress an autosomal dominant disease- related gene. In yet further embodiment, two or more sgRNA(s) are used to "Chop", remove, or suppress an autosomal dominant disease- related gene.

**[0114]** In one embodiment, the DNA digesting agent can be a site-specific nuclease. In another embodiment, the site-specific nuclease may be a Cas-family nuclease. In a more specific embodiment, the Cas nuclease may be a Cas9 nuclease.

**[0115]** In one embodiment, Cas protein may be a functional derivative of a naturally occurring Cas protein.

**[0116]** In addition to well characterized CRISPR-Cas system, a new CRISPR enzyme, called Cpf1 (Cas protein 1 of PreFran subtype) has recently been described (Zetsche et al. Cell. pii: S0092-8674(15)01200-3. doi: 10.1016/j.cell.2015.09.038 (2015)). Cpf1 is a single RNA-guided endonuclease that lacks tracrRNA, and utilizes a T-rich protospacer-adjacent motif. The authors demonstrated that Cpf1 mediates strong DNA interference with characteristics distinct from those of Cas9. Thus, in one embodiment of the present invention, CRISPR-Cpf1 system can be used to cleave a desired region within the targeted gene.

**[0117]** In further embodiment, the DNA digesting agent is a transcription activator-like effector nuclease (TALEN). TALENs are composed of a TAL effector domain that binds to a specific nucleotide sequence and an endonuclease domain that catalyzes a double strand break at the target site (PCT Patent Publication No. WO2011072246; Miller et al., Nat. Biotechnol. 29, 143-148 (2011); Cermak et al., Nucleic Acid Res. 39, e82 (2011)). Sequence-specific endonucleases may be modular in nature, and DNA binding specificity is obtained by arranging one or more modules. Bibikova et al., Mol. Cell. Biol. 21, 289-297 (2001). Boch et al., Science 326, 1509-1512 (2009).

**[0118]** ZFNs can be composed of two or more (e.g., 2 - 8, 3 - 6, 6 - 8, or more) sequence-specific DNA binding domains (e.g., zinc finger domains) fused to an effector endonuclease domain (e.g., the FokI endonuclease). Porteus et al., Nat. Biotechnol. 23, 967-973 (2005). Kim et al. (2007) Hybrid restriction enzymes: Zinc finger fusions to Fok I cleavage domain, Proceedings of the National Academy of Sciences of USA, 93: 1156-1160. U.S. Patent No. 6,824,978. PCT Publication Nos. WO1995/09233 and WO1994018313.

**[0119]** In one embodiment, the DNA digesting agent is a site-specific nuclease of the group or selected from the group consisting of omega, zinc finger, TALE, and CRISPR/Cas.

**[0120]** The sequence-specific endonuclease of the methods and compositions described here can

be engineered, chimeric, or isolated from an organism. Endonucleases can be engineered to recognize a specific DNA sequence, by, e.g., mutagenesis. Seligman et al. (2002) Mutations altering the cleavage specificity of a homing endonuclease, *Nucleic Acids Research* 30: 3870-3879. Combinatorial assembly is a method where protein subunits form different enzymes can be associated or fused. Arnould et al. (2006) Engineering of large numbers of highly specific homing endonucleases that induce recombination to novel DNA targets, *Journal of Molecular Biology* 355: 443-458. In certain embodiments, these two approaches, mutagenesis and combinatorial assembly, can be combined to produce an engineered endonuclease with desired DNA recognition sequence.

**[0121]** The sequence-specific nuclease can be introduced into the cell in the form of a protein or in the form of a nucleic acid encoding the sequence-specific nuclease, such as an mRNA or a cDNA. Nucleic acids can be delivered as part of a larger construct, such as a plasmid or viral vector, or directly, e.g., by electroporation, lipid vesicles, viral transporters, microinjection, and biolistics. Similarly, the construct containing the one or more transgenes can be delivered by any method appropriate for introducing nucleic acids into a cell.

**[0122]** Single guide RNA(s) used in the methods of the present disclosure can be designed so that they direct binding of the Cas-sgRNA complexes to pre-determined cleavage sites in a genome. In one embodiment, the cleavage sites may be chosen so as to release a fragment or sequence that contains a region of autosomal dominant disease-related gene. In further embodiment, the cleavage sites may be chosen so as to release a fragment or sequence that contains a region of *RHO*.

**[0123]** For Cas family enzyme (such as Cas9) to successfully bind to DNA, the target sequence in the genomic DNA should be complementary to the sgRNA sequence and must be immediately followed by the correct protospacer adjacent motif or "PAM" sequence. "Complementarity" refers to the ability of a nucleic acid to form hydrogen bond(s) with another nucleic acid sequence by either traditional Watson-Crick or other non-traditional types. A percent complementarity indicates the percentage of residues in a nucleic acid molecule, which can form hydrogen bonds (e.g., Watson-Crick base pairing) with a second nucleic acid sequence. Full complementarity is not necessarily required, provided there is sufficient complementarity to cause hybridization and promote formation of a CRISPR complex. A target sequence may comprise any polynucleotide, such as DNA or RNA polynucleotides. The Cas9 protein can tolerate mismatches distal from the PAM, however, mismatches within the 12 base pairs (bps) of sequence next to the PAM sequence can dramatically decrease the targeting efficiency. The PAM sequence is present in the DNA target sequence but not in the sgRNA sequence. Any DNA sequence with the correct target sequence followed by the PAM sequence will be bound by Cas9. The PAM sequence varies by the species of the bacteria from which Cas9 was derived. The most widely used CRISPR system is derived from *S. pyogenes* and the PAM sequence is NGG located on the immediate 3' end of the sgRNA recognition sequence. The PAM sequences of CRISPR systems from exemplary bacterial species include: *Streptococcus pyogenes* (NGG), *Neisseria meningitidis* (NNNNGATT), *Streptococcus thermophilus* (NNAGAA) and *Treponema denticola* (NAAAAC).

**[0124]** sgRNA(s) used in the present disclosure can be between about 5 and 100 nucleotides long, or longer (e.g., 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28,

29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, or 100 nucleotides in length, or longer). In one embodiment, sgRNA(s) can be between about 15 and about 30 nucleotides in length (e.g., about 15-29, 15-26, 15-25; 16-30, 16-29, 16-26, 16-25; or about 18-30, 18-29, 18-26, or 18-25 nucleotides in length).

**[0125]** To facilitate sgRNA design, many computational tools have been developed (See Prykhozhij et al. (PLoS ONE, 10(3): (2015)); Zhu et al. (PLoS ONE, 9(9) (2014)); Xiao et al. (Bioinformatics, Jan 21 (2014)); Heigwer et al. (Nat Methods, 11(2): 122-123 (2014)). Methods and tools for guide RNA design are discussed by Zhu (Frontiers in Biology, 10 (4) pp 289-296 (2015)). Additionally, there is a publically available software tool that can be used to facilitate the design of sgRNA(s) (<http://www.genscript.com/gRNA-design-tool.html>).

**[0126]** A modified autosomal dominant disease-related gene or fragment sequence is a donor sequence that has been codon modified to be unrecognizable by sgRNA(s) used for targeting or recognition of the mutated autosomal dominant disease-related gene and resistant to sgRNA targeting. Such modified autosomal dominant disease-related gene sequence is a donor sequence encoding at least a functional fragment of the protein lacking or deficient in the subject with autosomal dominant disease.

**[0127]** As previously mentioned, the codon-modified cDNA (donor-template) may be modified in such a way as to render it unrecognizable by the sgRNA(s) used to target either mutant and wildtype disease-related gene(s). To achieve this, mutations need to be introduced into a donor-template gene or fragment to render donor-template gene or fragment unrecognizable by sgRNA(s) and consequently resistant to degradation by Cas enzyme (such as Cas9 nuclease) which has been introduced in cells. The donor-template gene may be modified by introducing a wobble base(s) into donor-template. Introduction of wobble base(s) in DNA results in a silent mutation, leaving the expression product of wt gene intact, but if nucleotide sequence has been sufficiently changed, it will render donor-template sequence unrecognizable by sgRNA(s) used to target either mutant and wt disease-related gene(s), ultimately resistant to Cas nuclease cleavage. The number of wobble bases that needs to be introduced into a donor-template may vary, but needs to be sufficient to prevent sgRNA hybridization and formation of a CRISPR complex.

**[0128]** In one embodiment, the donor template sequence may be delivered using the same gene transfer system as used to deliver the Cas nuclease (included on the same vector) or may be delivered using a different delivery system. In another embodiment, the donor template sequence may be delivered using the same transfer system as used to deliver sgRNA(s). In specific embodiments, the donor is delivered using a viral vector (e.g., AAV).

**[0129]** In one embodiment, the present disclosure comprises integration of codon-modified autosomal dominant disease-related gene sequence (donor template sequence) into the endogenous autosomal disease-related gene.

**[0130]** In another embodiment, the donor sequence or modified autosomal dominant disease-related gene sequence is integrated into endogenous gene by homologous recombination (HR).

**[0131]** In further embodiments, the donor sequence or modified autosomal dominant disease-related gene sequence is flanked by an upstream and a downstream homology arm. The homology arms, which flank the donor sequence or modified autosomal dominant disease-related gene sequence, correspond to regions within the targeted locus of autosomal dominant disease-related gene. For example, the corresponding regions within the targeted locus are referred to herein as "target sites". Thus, in one example, a vector that carries a donor or modified autosomal dominant disease-related gene sequence can comprise a donor or modified autosomal dominant disease-related gene sequence flanked by a first and a second homology arm.

**[0132]** A homology arm of the vector that carries a donor or modified autosomal dominant disease-related gene sequence can be of any length that is sufficient to promote a homologous recombination event with a corresponding target site, including for example, 50-100 base pairs, 100-1000 base pairs or at least 5-10, 5-15, 5-20, 5-25, 5-30, 5-35, 5-40, 5-45, 5-50, 5-55, 5-60, 5-65, 5-70, 5-75, 5-80, 5-85, 5-90, 5-95, 5-100, 100-200, or 200-300 base pairs in length or greater.

**[0133]** In one embodiment, the donor template is delivered as a double-stranded DNA. Under such circumstances, the homologous arm may comprise 15-4000 base pairs of each arm. In other embodiments, the donor template is delivered as a single-stranded DNA format. Under such circumstances, the homologous arm may comprise 8-1000 bps of each arm.

**[0134]** A homology arm and a target site "correspond" or are "corresponding" to one another when the two regions share a sufficient level of sequence identity to one another to act as substrates for a homologous recombination reaction. By "homology" is meant DNA sequences that are either identical or share sequence identity to a corresponding sequence. The sequence identity between a given target site and the corresponding homology arm found on the vector that carries a donor or modified autosomal dominant disease-related gene sequence can be any degree of sequence identity that allows for homologous recombination to occur. For example, the amount of sequence identity shared by the homology arm of the vector that carries a donor or modified autosomal dominant disease-related gene sequence (or a fragment thereof) and the target site (or a fragment thereof) should be 100% sequence identity, except the codon-modified region, such that the sequences undergo homologous recombination. Less than 100% sequence identity may be tolerated, provided that the Cas enzyme (Cas 9) cuts only the patient DNA and not the donor template or the patient DNA which is repaired/replaced by the donor template.

**[0135]** Alternatively, donor template (whether codon-modified or not) of a gene of interest or fragment is not integrated into the endogenous disease-related gene. Donor-template may be packaged into an extrachromosomal, or episomal vector (such as AAV vector), which persists in the nucleus in an extrachromosomal state, and offers donor-template delivery and expression without integration into the host genome. Use of extrachromosomal gene vector technologies has been discussed in detail by Wade-Martins R (Methods Mol Biol. 2011;738:1-17).

**[0136]** The nucleases, polynucleotides encoding these nucleases, donor polynucleotides and compositions comprising the proteins and/or polynucleotides described in the present disclosure can be delivered by any suitable means. In certain embodiments, the nucleases and/or donors are delivered *in vivo*. In other embodiments, the nucleases and/or donors are delivered to isolated cells (e.g., autologous iPS cells) for the provision of modified cells useful in *in vivo* delivery to patients

afflicted with ocular autosomal dominant disease.

**[0137]** An alternative to injection of viral particles encoding CRISPR components described in the present disclosure (including sgRNA(s), codon-modified donor template gene or fragment sequences, and Cas family nuclease), cell replacement therapy can be used to prevent, correct or treat diseases, where the methods of the present disclosure are applied to isolated patient's cells (*ex vivo*), which is then followed by the injection of "corrected" cells back into the patient.

**[0138]** For the treatment of ocular diseases, patient's iPS cells can be isolated and differentiated into retinal pigment epithelium RPE cells *ex vivo*. RPE cells characterized by the mutation in autosomal dominant disease-related gene may then be manipulated using methods of the present disclosure in a manner that results in the deletion of autosomal dominant disease-related gene, and expression of a corrected autosomal dominant disease-related gene.

**[0139]** Thus, the present disclosure provides methods for correcting autosomal dominant ocular disease in a subject, wherein the method results in functional recovery of the autosomal dominant ocular disease-related gene, comprising administering to the subject a therapeutically effective amount of autologous differentiated retinal pigment RPE cells expressing a corrected autosomal dominant ocular disease-related gene. Administration of the pharmaceutical preparations comprising autologous RPE cells that express a corrected autosomal dominant ocular disease-related gene may be effective to reduce the severity of symptoms and/or to prevent further deterioration in the patient's condition. Such administration may be effective to fully restore any vision loss or other symptoms.

**[0140]** For example, patient fibroblast cells can be collected from the skin biopsy and transformed into iPS cells. Dimos JT et al. (2008) Induced pluripotent stem cells generated from patients with ALS can be differentiated into motor neurons. *Science* 321: 1218-1221, *Nature Reviews Neurology* 4, 582-583 (November 2008). Luo et al., Generation of induced pluripotent stem cells from skin fibroblasts of a patient with olivopontocerebellar atrophy, *Tohoku J. Exp. Med.* 2012, 226(2): 151-9. The CRISPR-mediated correction can be done at this stage. The corrected cell clone can be screened and selected by RFLP assay. The corrected cell clone is then differentiated into RPE cells and tested for its RPE-specific markers (Bestrophin1, RPE65, Cellular Retinaldehyde-binding Protein, and MFRP). Well-differentiated RPE cells can be transplanted autologously back to the donor patient.

**[0141]** The well-differentiated autologous RPE cells described in the present disclosure may be formulated with a pharmaceutically acceptable carrier. For example, autologous RPE cells can be administered alone or as a component of a pharmaceutical formulation. The autologous RPE cells of the present disclosure can be administered in combination with one or more pharmaceutically acceptable sterile isotonic aqueous or nonaqueous solutions (e.g., balanced salt solution (BSS)), dispersions, suspensions or emulsions, or sterile powders which may be reconstituted into sterile injectable solutions or dispersions just prior to use, which may contain antioxidants, buffers, bacteriostats, solutes or suspending or thickening agents.

**[0142]** The autologous RPE cells of the present disclosure may be delivered in a pharmaceutically acceptable ophthalmic formulation by intraocular injection. Concentrations for injections may be at

any amount that is effective and nontoxic. The pharmaceutical preparations of autologous RPE cells of the present disclosure for treatment of a patient may be formulated at doses of at least about  $10^4$  cells/mL. The RPE cell preparations for treatment of a patient can be formulated at doses of at least about  $10^3$ ,  $10^4$ ,  $10^5$ ,  $10^6$ ,  $10^7$ ,  $10^8$ ,  $10^9$ , or  $10^{10}$  RPE cells/mL.

**[0143]** Subjects, which may be treated according to the present invention, include all animals which may benefit from the present invention. Such subjects include mammals, preferably humans (infants, children, adolescents and/or adults), but can also be an animal such as dogs and cats, farm animals such as cows, pigs, sheep, horses, goats and the like, and laboratory animals (e.g., rats, mice, guinea pigs, and the like).

## **EXAMPLES**

### **Surveyor assay**

**[0144]** Surveyor mutation detection assay provides a simple and robust method to detect mutations and polymorphisms in DNA mixture. The key component of the kit is Surveyor Nuclease, a member of the CEL family of mismatch-specific nucleases derived from celery. Surveyor Nuclease recognizes and cleaves mismatches due to the presence of single nucleotide polymorphisms (SNPs) or small insertions or deletions.

**[0145]** Surveyor nuclease cleaves with high specificity at the 3' side of any mismatch site in both DNA strands, including all base substitutions and insertion/deletions up to at least 12 nucleotides. Surveyor nuclease technology involves four steps: (i) PCR to amplify target DNA from the cell or tissue samples underwent Cas9 nuclease-mediated cleavage (here we expect to see a nonhomogeneous or mosaic pattern of nuclease treatment on cells, some cells got cuts, some cells don't); (ii) hybridization to form heteroduplexes between affected and unaffected DNA (Because the affected DNA sequence will be different from the unaffected, a bulge structure resulted from the mismatch can form after denature and renature); (iii) treatment of annealed DNA with Surveyor nuclease to cleave heteroduplexes (cut the bulges); and (iv) analysis of digested DNA products using the detection/separation platform of choice, for instance, agarose gel electrophoresis. The Cas9 nuclease-mediated cleavage efficacy can be estimated by the ratio of Surveyor nuclease-digested over undigested DNA. The technology is highly sensitive, detecting rare mutants present at as low as 1 in 32 copies. Surveyor mutation assay kits are commercially available from Integrated DNA Technologies (IDT), Coraville, IA.

### **RFLP analysis**

**[0146]** Restriction fragment length polymorphism (RFLP) analysis is a technique well-known to those skilled in the art. RFLP exploits variations in homologous DNA sequences. The basic technique for the detecting RFLPs involves fragmenting a sample of DNA by a restriction enzyme, which can recognize and cut DNA wherever a specific short sequence occurs, in a process known

as a restriction digest. The resulting DNA fragments are then separated by length on agarose gel electrophoresis for analysis. For the detection of donor-template replacement (also known as gene-correction), one or multiple kinds of additional restriction enzyme sites are introduced into the donor template by codon-modification, without affecting the overall length. After using PCR to amplify the target DNA sequence from tissue samples, the PCR amplicon can be evaluated by the aforementioned restriction enzyme(s) for the detection of the samples that underwent gene correction.

**[0147]** The following examples of specific aspects for carrying out the present invention are offered for illustrative purposes only, and are not intended to limit the scope of the present invention in any way.

### **Example 1**

#### ***ChopStick AAV Gene Therapy Strategy Evaluation***

**[0148]** The present Example outlines the strategy behind ChopStick AAV gene therapy. The approach is based on using a gene-editing enzyme with one or more unique single guide RNA (sgRNA) sequence that target both mutant and wild type forms of rhodopsin for destruction. This initial step is then followed by supplying a wild-type codon modified rhodopsin cDNA to the cells. A significant advantage of this system is that the codon modified rhodopsin cDNA is not recognizable by the sgRNA(s) and thus is resistant to the cleavage by the nuclease.

**[0149]** As shown in Figure 1, the "ChopStick" system described here is packaged into two recombinant AAV vectors (Fig. 1A). The first vector carries the polynucleotide sequence encoding the Cas9 enzyme (SEQ ID NO: 17), which is able to "chop" the mutant and native rhodopsin genes, while the second vector contains a polynucleotide encoding the codon-modified human rhodopsin to "stick" the normal rhodopsin back into the patient. The codon-modified engineered human rhodopsin sequence, which in this example is driven by the CBh promoter (SEQ ID NO: 10), is resistant to destruction by the gene-editing enzyme (Cas9 in this instance), and allows for the rescue of patient's phenotype. In addition to carrying a codon-modified engineered human rhodopsin sequence, the second vector carries a two single guide RNAs (sgRNA1 and sgRNA2) which act as a guide to define the target site to introduce DNA double-stranded break and thus acts as a homing device for directing the Cas9 nuclease. Each pair of recombinant AAV vectors can be used to target rhodopsin genes. The codon-modified sequence is shown in Figure 1B section II. Each sgRNA targeting site comprises four mismatches which are underlined.

**[0150]** AAV has a packaging capacity of 4.5~4.9Kb. Since the coding sequence of spCas9 is ~ 4.2Kb and the two inverted terminal repeat (ITRs) of AAV is ~0.3 Kb, there is about 0.4 Kb of space for promoter and poly-adenine termination signal. The inventors of the present disclosure used a 173 bp short CMV promoter and a 50 bp synthetic poly-adenine signal to construct the Cas AAV vector. The detailed sequence is listed and all of the components of the recombinant AAV vector are shown below: Underline and bold: ITR (SEQ ID NO: 13); Underlined: short CMV promoter;



CTGACCCCTGACACTGTTTGAGGACAGAGAGATGATCGAGGAACGGCTGAAAAACCTA  
TGCCACCTGTTCGACGACAAAAGTGATGAAGCAGCTGAAGCGGCGGAGATACACCG  
GCTGGGGCAGGCTGAGCCGGAAGCTGATCAACGGCATCCGGGACAAGCAGTCCGGC  
AAGACAATCCTGGATTTCCTGAAGTCCGACGGCTTCGCCAACAGAACTTCATGCAG  
CTGATCCACGACGACAGCCTGACCTTTAAAGAGGACATCCAGAAAGCCCAGGTGTC  
CGGCCAGGGCGATAGCCTGCACGAGCACATTGCCAATCTGGCCGGCAGCCCCGCCA  
TTAAGAAGGGCATCCTGCAGACAGTGAAGGTGGTGGACGAGCTCGTGAAAGTGATG  
GGCCGGCACAAGCCCGAGAACATCGTGATCGAAATGGCCAGAGAGAACCAGACCA  
CCCAGAAGGGACAGAAGAACAGCCGCGAGAGAATGAAGCGGATCGAAGAGGGCAT  
CAAAGAGCTGGGCAGCCAGATCCTGAAAGAACACCCCGTGGAACACCCAGCTGC  
AGAACGAGAAGCTGTACCTGTACTACCTGCAGAATGGGCGGGATATGTACGTGGAC  
CAGGAACGGACATCAACCGGCTGTCCGACTACGATGTGGACCATATCGTGCCTCA  
GAGCTTTCTGAAGGACGACTCCATCGACAACAAGGTGCTGACCAGAAGCGACAAGA  
ACCGGGGCAAGAGCGACAACGTGCCCTCCGAAGAGGTTCGTGAAGAAGATGAAGAA  
CTACTGGCGGCAGCTGCTGAACGCCAAGCTGATTACCCAGAGAAAAGTTCGACAATC  
TGACCAAGGCCGAGAGAGGGCGGCCTGAGCGAACTGGATAAAGCCGGCTTCATCAAG  
AGACAGCTGGTGGAAACCCGGCAGATCACAAGCACGTGGCACAGATCCTGGACTC  
CCGGATGAACACTAAGTACGACGAGAATGACAAGCTGATCCGGGAAGTGAAAGTGA  
TCACCCTGAAGTCCAAGCTGGTGTCCGATTTCCGGAAGGATTTCCAGTTTTACAAAG  
TGCGCGAGATCAACAACACTACCACCACGCCACGACGCCTACCTGAACGCCGTCTGTG  
GGAACCGCCCTGATCAAAAAGTACCCTAAGCTGGAAAGCGAGTTCGTGTACGGCGA  
CTACAAGGTGTACGACGTGCGGAAGATGATCGCCAAGAGCGAGCAGGAAATCGGCA  
AGGCTACCGCCAAGTACTTCTTCTACAGCAACATCATGAACTTTTTCAAGACCGAGA  
TTACCCTGGCCAACGGCGAGATCCGGAAGCGGCCTCTGATCGAGACAAACGGCGAA  
ACCGGGGAGATCGTGTGGGATAAAGGGCCGGGATTTTGCCACCGTGCGGAAAGTGCT  
GAGCATGCCCAAGTGAATATCGTGAAAAAGACCGAGGTGCAGACAGGCGGCTTCA  
GCAAAGAGTCTATCCTGCCCAAGAGGAACAGCGATAAGCTGATCGCCAGAAAAGAAG  
GACTGGGACCCTAAGAAGTACGGCGGCTTCGACAGCCCCACCGTGGCCTATTCTGTG  
CTGGTGGTGGCCAAAGTGGAAGGGCAAGTCCAAGAACTGAAGAGTGTGAAAG  
AGCTGCTGGGGATCACCATCATGGAAAGAAGCAGCTTCGAGAAGAATCCCATCGAC  
TTTCTGGAAGCCAAGGGCTACAAAGAAGTGAAAAAGGACCTGATCATCAAGCTGCC  
TAAGTACTCCCTGTTTCGAGCTGGAAAACGGCCGGAAGAGAATGCTGGCCTCTGCCG  
GCGAACTGCAGAAGGGAAACGAACTGGCCCTGCCCTCCAAATATGIGAACCTTCTG  
TACCTGGCCAGCCACTATGAGAAGCTGAAGGGCTCCCCGAGGATAATGAGCAGAA  
ACAGCTGTTTGTGGAACAGCACAAGCACTACCTGGACGAGATCATCGAGCAGATCA  
GCGAGTTCTCCAAGAGAGTGATCCTGGCCGACGCTAATCTGGACAAAGTGCTGTCCG  
CCTACAACAAGCACCGGGATAAGCCCATCAGAGAGCAGGCCGAGAATATCATCCAC  
CTGTTTACCCTGACCAATCTGGGAGCCCCTGCCGCTTCAAGTACTTTGACACCACC  
ATCGACCGGAAGAGGTACACCAGCACCAAAGAGGTGCTGGACGCCACCCTGATCCA  
CCAGAGCATCACCGCCTGTACGAGACACGGATCGACCTGTCTCAGCTGGGAGGGC  
ACAAAAGGCCGGCGGCCACGAAAAAGGCCGGCCAGGCCAAAAAAGAAAAAGTAAgTc



lane 4). Using one sgRNA (lanes 2 and 3) at a time does not result in change in size. Approximately 30% of the genomic DNA underwent non-homologous end joining (NHEJ) by one sgRNA, and up to 80% was edited (deletion and NHEJ) when two sgRNAs were used. Equal amounts of plasmid DNA (1  $\mu\text{g}$  /  $1 \times 10^5$  293FT cells) were used in each group. These findings indicate that applying two sgRNAs can destroy, remove, or degrade endogenous human *RHO* sequence more efficiently.

**[0152]** The inventors further tested if the "Chop" can decrease wt *RHO* gene expression. In this experiment, a bicistronic construct was transfected into HEK293 cells to express wt *RHO* cDNA (SEQ ID NO: 8) and EGFP (Fig. 3A). Both *RHO* cDNA and EGFP expression (SEQ ID NO: 21) were driven by a CMV promoter (SEQ ID NO: 20) independently and simultaneously, so that EGFP expression can be used as an internal control in western blot, which normalizes the difference in transfection efficiency and protein loading. Figure 3A also illustrates the target sites of sgRNA1 (SEQ ID NO: 1) and sgRNA2 (SEQ ID NO: 2) on this *RHO* expression vector. When HEK293FT cells were co-transfected with *RHO* expression vector and another vector expressing Cas9 components (pX459) carrying sgRNA1 and sgRNA2, the *RHO* protein expression level was much lower (Fig. 3B). The sg3 group is a non-specific control sgRNA. These result were further normalized with internal control, which is shown in Figure 3C. These findings indicate that, applying two sgRNAs together lowers *RHO* expression about 70%, while using single sgRNA only reduced 0-30% compared to the control group (sg3). Together, these results indicate that "Chop" strategy can significantly abolish or inactivate wt *RHO* protein expression.

## **Example 2**

### ***CRISPR/Cas9-Induced***

#### ***Gene Editing in a Mouse Model of Retinitis Pigmentosa Delays Disease Progression***

**[0153]** Next, the inventors verified the feasibility and efficacy of the CRISPR/Cas9 endonuclease system as a gene-editing treatment modality in a mouse model of RP with the dominant D190N rhodopsin mutation. In these experiments, two AAV8 vectors containing the Cas9 coding sequence and the sgRNA (SEQ ID NO: 4)/donor template marked with an *Afl*III restriction site were used. Insertion of *Afl*III restriction site allows for the identification of cells that have undergone homologous recombination (Figs. 4A-4C). Briefly, heterozygous  $\text{Rho}^{\text{D190N}/+}$  was transduced into the right eye before post-natal day 5 with above described recombinant AAV8 vectors. The sgRNA targeting frequency and recombination of donor template (SEQ ID # 23) were verified by TIDE indel tracking tool (Brinkman et al. Nucleic Acid Res. 2014 Dec 16, 42(22): e168) and *Afl*III enzyme digestion (Fig. 4). About 50% of cells underwent NHEJ (mostly are 1 bp insertion), and about 10% of cells incorporated donor template successfully. Structural preservation was assessed by H&E staining, and retinal function rescue was assessed by electroretinography (ERG) at 3 months of age (Figs. 5A and 5B). In the  $\text{Rho}^{\text{D190N}/+}$  3 month old mice, treated eyes showed greater photoreceptor survival than did eyes that did not receive AAV injection (Fig. 5A). Furthermore, retinal function as

measured by ERG was also increased in treated eyes compared with control eyes at 3 months of age (Fig. 5B). Collectively, these results demonstrate that CRISPR-Cas9 gene editing described in the present disclosure can be used to *in vivo* correct the phenotype of RP. The codon-modified donor sequence is shown as follows: Underline and bold: homologous arm; **Bold**: Afill site; Underlined: 5 codon-modified wobble nucleotides;

tccttaaccaccgaaggcagggcagcaggctagtgagcagagctgcctggcaagtggcagggagcttaagaatcgtccaag  
ggcggagaccagtaagtctcattaggtgatggggcagcaggtaaaagccattcatGcttatgtccagctggcggtgtttctette  
ctgtttatcacccttgcctgaccatcaggtacatccctgagggcatgcaatgttcatgcgggattgactattatacccttaagccgg  
aggtcaacaacgaatcctttgcatctacatgttcgtggctccactcaccattcctatgatcgtcatcttcttctctatgggcagctggt  
cttcacagtcaaggaggtatgagcaggg.

### Example 3

#### **CRISPR-Mediated Humanized Exon 1 at the Mouse Rho Locus**

**[0154]** The inventors of the present disclosure next tested the ability to replace mouse Rho locus with wild-type (wt) (SEQ ID NO: 24) or mutant human (h) (SEQ ID NO: 25) *RHO* exon 1 in mouse embryonic stem (ES) cells (Fig. 6). Briefly, ES cells were co-transfected (via electroporation) with the Cas9 expression vector carrying a *Rho* exon 1-specific sgRNA (sgRNA-Rho Exon 1, SEQ ID NO: 5) targeting mouse *Rho* exon 1) and a targeting vector carrying with human RHO donor template, which contained a sequence of hRHO exon 1 flanked with ~750 bp homologous arm on each side (Fig. 6A). Human *RHO* donor template is expected to replace mouse exon 1 and confer resistance to sgRNA-Rho Exon 1. Seven days after electroporation, ES clones were picked and DNA was extracted and amplified with screening primers. Two out of 96 clones were detected with replacement of human exon 1 by RFLP analysis (Fig. 6B). As shown in Fig. 6C, sequence electropherograms of amplicons show perfect fused human and mouse sequence of one targeted ES clone (lane 2, Fig 6B). The correct targeted clones can be further used to produce the humanized RHO exon 1 mouse model. This patientspecific humanized mouse system enables the inventors to test various sgRNAs that may be used for targeting human genomic sequence for ChopStick strategy *in vivo*. The advantages of using these mouse models also enables the validation of the "ChopStick" efficacy and safety via functional evaluation methods like visual function, imaging of rescued tissue in live animals for long term observations. The sgRNA sequence is listed above and the donor template sequence is listed below: Underline and bold: homologous arm; UPPERCASE: human *RHO* exon 1

atgctcaactgaataacctggcagcctgctccctcatgcagggaccacgtctctgctgcacccagcaggccatcccgctccatagcc  
catggtcatccctccctggacaggaatgtgtctctcccccggctgagcttctcaagctagaagcactccgaacagggttatggg  
cgctctccatctcccaagtggctggtatgaaatgttaatgtacatgtgagtgaaacaaattccaattgaacgcaacaatagtta  
tcgagccctgagccgggggggggggtgtgagactggaggcagtgacggagctgacggcacacacagctcagatctgtca  
agtgagccattgtcagggcttgggactggataagtcaggggtctcctgggaagagatgggataggtgagttcaggaggagac  
attgtcaactggagccatgtggagaagtgaatttagggcccaaggttcagctgcagcctgaggccaccagactgacatgggga  
ggaattccagaggactctggggcagacaagatgagacaccttctcttcttacctaaagggcctccaccggatgtcacttggcc  
cctctgcaagccaattaggccccggtggcagcagtgaggatfagcgttagtatgatactcgggatgctgaatcagcctctgctta

gggagagaaggctcactttataagggctctgggggggctcagtcctggagttgctgtgggagcGAGTCATCCAGCTG  
 GAGCCCTGAGTGAGTGGCTGAGCTCAGGCCTTCGCAGCATTCTTGGGTGGGAGCAGCCAC  
 GGGTCAGCCACAAGGGCCACAGCCATGAATGGCACAGAAGGCCCTAACTTCTACGT  
 GCCCTTCTCCAATGCGACGGGTGTGGTACGCAGCCCCTTCGAGTACCCACAGTACTA  
 CCTGGCTGAGCCATGGCAGTTCTCCATGCTGGCCGCCTACATGTTTCTGCTGATCGTG  
 CTGGGCTTCCCCATCAACTTCCTCACGCTCTACGTACCCGTCCAGCACAGAAGCTG  
 CGCACGCCTCTCAACTACATCCTGCTCAACCTAGCCGTGGCTGACCTCTTCATGGTC  
 CTAGGTGGCTTACCAGCACCCCTCTACACCTCTCTGCATGGATACTTCGTCTTCGGGC  
 CCACAGGATGCAATTTGGAGGGCTTCTTTGCCACCCTGGGCGgtatgagcagagagactggggc  
gggggggtagcatgggagccaagggccacgaaaagggcctgggagggctcagccttacttgagctctttaattggtctc  
taaagggccagcttattcattggcaaacactgtgacctgagctaggctgctgtgagagcaggcacggaacattcatctatctcat  
ctgagcaatgcaagaacatgggttcagagagggccaaggactcaccgaggagtcacagagtggtgggggtgctctgaggcag  
ctgagctggggcacacacagactgagcaccaggagtgagctctagctttttttctatgtcttttctaaaagacacataggtta  
ggactgtccctgggtccagtaagaactggttcagtaaaactgtacatctcactgctggccagccctgctcagcttccaccagagtcg  
tgcactacacacccggcatctcaaggattcattctctatcttctctatctttggagtgaggecacagtctcagctagtcagtcagact  
ggccttaattctgcagctgaggatgacttaaaactgtcactctctctgcccagcctctcaagtctctgacacagccagccacca  
ctatgctacgcccagggtgttccaacattttctctcccttaactggaaggctcaatgaggctcttctgagaagcaacagagcc

#### Example 4

##### *Repair of Mouse Pde6a D670G Allele in Stem Cells*

**[0155]** Mutations in genes encoding subunits of the rod-specific enzyme, cyclic guanosine monophosphate (cGMP) phosphodiesterase 6 (PDE6A and PDE6B), are responsible for approximately 72,000 cases of RP worldwide each year, making therapeutic modeling highly relevant to developing mechanisms based therapies. In the present Example, the inventors used the CRISPR/ Cas9 gene editing system to correct photoreceptor gene mutations in mouse ES cells.

**[0156]** For these studies, the inventors used ES cells isolated from Pde6a<sup>D670G/D670G</sup> mouse model (Wert KJ et al. Hum Mol Genet. 2013 Feb 1;22(3):558-67; Wert KJ et al. J Vis Exp. 2012 Nov 25; (69)). As shown in Fig. 7A, a donor construct (SEQ ID NO: 26) used in this Example contains two modifications: 1) a Pde6a-codon modification which creates an additional SphI site upstream from the D670G codon, where SphI enzyme-digestion can identify ES cells that underwent CRISPR-mediated homologous recombination; and 2) eight wobble base pairs were introduced, which make the donor template unrecognizable to sgRNA (SEQ ID NO: 6) and thus resistant to Cas9, and which resulted in the change of the mutant amino acid sequence to that of the wt amino acid sequence. Thus, upon Cas9 cutting and homologous recombination, the endogenous mutant allele is replaced (i.e., repaired). In Figure 7A, triangle indicates the sgRNA target site while the two arrows represent the primer pairs used for PCR amplification. As shown in Figure 7B, amplicons generated from recombined cells were 303 bp and 402 bp while the unedited amplicon is 705 bp. Moreover, using

direct sequencing of genomic DNA from a target clone, the inventors confirmed predicted replacement of the D670G exon with donor template (Fig. 7C). This is an example where there is no sgRNA target site on the mutation site, but researchers can still design sgRNA nearby and successfully replace mutant allele through homologous recombination.

**[0157]** Thus, in this Example, the inventors verified the ability of the CRISPR/Cas9 system to edit the mouse *PDE6a* locus and rescue photoreceptors.

#### Example 5

##### ***CRISPR/Cas9-Induced In Vivo Gene Editing in *Pde6a*<sup>D670G</sup> /*Pde6a*<sup>D670G</sup> Mouse Model***

**[0158]** The inventors have verified the use of CRISPR/Cas9 system to edit the mouse *Rho* locus and rescue photoreceptors (Example 2). Furthermore, as shown in Example 4, the inventors were able to repair mouse *Pde6a* D670G allele in ES cells. Next, the inventors will perform *in vivo* experiments, where post-natal day (P) 5 *Pde6a*<sup>D670G</sup> /*Pde6a*<sup>D670G</sup> mice will receive subretinal transductions of both recombinant AAV8-Cas9 and AAV8-sgRNA with the codonoptimized Cas9 resistant donor DNA (validated in Example 4) into one eye. In control animals, one eye will be transduced with an empty AAV8 vector or AAV8-Cas9 as negative control.

**[0159]** The inventors will next perform quantitative validation of recombination and correction of one of the *Pde6a*<sup>D670G</sup> allele in homozygous mutant. Briefly, one month after injection (before degeneration onset), retinas will be dissected, DNA isolated, PCR performed, and SphI restriction site verified (using RFLP). PCR samples will be run in triplicate. At 3 weeks of age, retinas from 3 mice will be collected, and *Pde6a* levels quantified by immunoblotting, as described.

**[0160]** To quantitatively assess photoreceptor function and survival, the inventors will perform quantitative AF of outer segment thinning on SD-OCT, ERG, and rod-cone density at 8, 16, and 24 weeks (n = 36 total animals) using previously published techniques (Woodruff et al. J Neurosci. 2008 Feb 27;28(9):2064-74; Janisch et al. Biochem Biophys Res Commun 390, 1149-1153 (2009); Tsang et al. Science. 1996 May 17;272(5264):1026-9; Davis et al. Invest Ophthalmol Vis Sci. 2008 Nov;49(11):5067-76.) All measurements will be performed on both eyes.

**[0161]** The inventors will also determine the efficacy of PDE function. As a key biochemical indicator of rescue, the inventors will measure whether total cGMP levels and PDE activity from light- and dark-adapted retinas are restored. Three additional sample right eyes, treated at P18, P21, P28, and P35, and control fellow left eyes of *Pde6a*<sup>D670G/D670G</sup> will be assayed. GUCY2E (guanylate cyclase) should remain stable for all experiments and will be determined as previously described (Tsang et al. Science. 1996 May 17; 272(5264): 1026-1029; Science. 1998 Oct 2;282(5386):117-21).

#### Example 6

***Efficacy and frequency of homologous recombination vs. non-homologous end joining (NHEJ) for editing PD6A in patient specific stem cells***

**[0162]** In this Example, the inventors will verify that the AAV2-Cas9 system can edit the human 6 *PDE6A* locus. In this aim,  $0.25 \times 10^6$  patient iPS in a 6-well matrigel coated plate (in NutriStem XF/FF Culture media, Stemgent, Cambridge) will be co-transduced with AAV2-Cas9 and AAV2 vector-donor template mix (MOI: 2000) to repair *PDE6A*. After 48 hours, iPS will be passaged with Accutase onto regular 10-cm matrigel coated culture dishes.

**[0163]** Next, 1000 *PDE6A*<sup>R102C</sup>/*PDE6A*<sup>S303C</sup> patient iPS clones will be picked, and DNA isolated for PCR; (Primers: forward: GCAGACTGCAAACTGCCAT; reverse: TGTCACCAGCCTTGTCTTGG). PCR products will be cut with *BsiWI* to identify clones that have undergone homologous recombination. After *BsiWI* digestion, the amplicon generated from iPS that underwent homologous recombination gives bands at 271 bp and 380 bp, compared to the parental sequence, which gives only one band at 651 bp.

**[0164]** To determine the percentage of clones that have undergone NHEJ (as opposed to those that underwent homology-directed repair), DNA will be analyzed from clones without the *BsiWI* site (i.e., not transduced, transduced off-target or NHEJ), and the frequency of the disruption of the *PDE6A* allele determined. DNA will be analyzed by SURVEYOR mismatch detection assay and positive DNA samples will be subjected to subcloned into plasmid vectors such as pCR™4Blunt-TOPO® vector and then send for Sanger sequencing. In addition, the assessment of off-targeting in iPS and live mice are prerequisites before application to humans. Off-target sites will be analyzed by full-genome sequencing using Illumina next-generation sequencing.

**[0165]** The inventors anticipate a much higher rate of homologous recombination mediated by AAV8 in photoreceptors *in vivo*, compared to patient iPS. This is because AAV8 introduces sgRNA into photoreceptors at a much higher frequency than transduction into iPS. AAV8 introduces into photoreceptors ~10,000 copies of the sgRNA and Cas9, as opposed to both lipofection and electroporation, which generally introduce a single-copy DNA into each cell.

**[0166]** The percentage of transduced clones that undergo NHEJ is likely to be higher than those undergoing homologous recombination-approximately 10% vs. 1%, respectively, with a 90-bp donor template in human- induced pluripotent stem cells.

**Example 7**

***Use of CRISPR system to replace mutant allele R345W in iPS cells isolated from Dooye Honeycomb patient***

**[0167]** Doyme Honeycomb retinal dystrophy (DHRD) is an inherited disease that affects the eyes and causes vision loss. It is characterized by small, round, white spots (drusen) that accumulate around the retinal pigment epithelium. Over time, drusen may grow and come together, creating a honeycomb pattern. It usually begins in early adulthood, but the age of onset varies. The degree of vision loss also varies. DHRD is caused by R345W mutations in the EFEMP1 gene, which are inherited in an autosomal dominant manner.

**[0168]** In this Example, the inventors used CRISPR components and a donor template (SEQ ID NO: SEQ ID 27) to correct the R345W mutation in the iPS cells derived from Doyme Honeycomb patient fibroblast (Figure 8A). The resulting iPS cells comprise wild type EFEMP1 sequence with codon-modification, which confers resistance to further cutting by the Cas9. These cells can be used for autologous transplantation after the differentiation into RPE cells for the cure of DHRD.

**[0169]** Briefly, the inventors collected the iPS derived from the DHRD patient. Cas9 protein and sgRNA-EFEMP1 (SEQ ID NO: 3) (Figure 8B) were mixed and co-transduced with donor template in the form of single strand oligodeoxynucleotide (ssODN) (sequence: tagttagtaaactcttgaccctacatctctacagatataaatgagtgtagaccacaaaCgaGtgcCgggaggatgaaatgtgttgaattatcatggcggctccgtgttatccacgaaatcctt) into iPS cells by nucleofection. The donor template is codon-modified to prevent repeating recognizing and cutting by the CRISPR components. The colony with corrected sequence was confirmed by RFLP assay with the additional ScrFI restriction site. The genotype of the iPS cell is further confirmed by sequencing (Figure 8C).

**[0170]** This experiment provides the evidence that the "Chop" strategy has potential to treat autosomal dominant diseases other than autosomal dominant retinitis pigmentosa.

**[0171]** Table 2 provides sgRNA sequences, donor-template modified sequences, and additional experiments used in the Examples of the present disclosure.

Table 2.

Sequence ID Number	Sequence	Species
SEQ ID NO: 1 (sgRNA1)	GGACGGTGACGTAGAGCGTG	<i>Homo sapiens</i>
SEQ ID NO: 2 (sgRNA2)	GACGAAGTATCCATGCAGAG	<i>Homo sapiens</i>
SEQ ID NO: 3 (sgRNA-EFEMP1)	TGAGACCACAAATGAATGCT	<i>Homo sapiens</i>
SEQ ID NO: 4 (sgRNA-D190N)	AACTACTACACACTCAAGCCTG	<i>Mus musculus</i>
SEQ ID NO: 5 (sgRNA-Rho Exon 1)	GTAGTACTGCGGCTGCTCGA	<i>Mus musculus</i>
SEQ ID NO: 6 (sgRNA- <i>Pde6a</i> <sup>D670G</sup> )	GCTCATGCTGCCGGCGATTC	<i>Mus musculus</i>
SEQ ID NO: 7 (sgRNA-YI)	GGTTTTGGACAATGGAACCG	<i>Drosophila melanogaster</i>
SEQ ID NO: 8 (wt)		<i>Homo</i>

Sequence ID Number	Sequence	Species
<i>RHO</i> cDNA)	<p>atgaatggcacagaaggccctaactctacgtgcccttctccaatgcga  cgggtgtggtacgcagcccctcagtagtaccacagtactacctggctg  agccatggcagttctccatgctggccgctacatgttctgctgatcgtg  ctgggcttcccatacaacttctcactgctctacgtcaccgtccagcacia  gaagctgcgcacgcctctcaactacatcctgctcaacctagccgtggct  gaccttctcaggtcctaggtggcttaccagcaccctctacacctctctg  catggafactcgtctcgggcccacaggatgcaattggaggcctcttt  gccaccctgggcgggaaattgccctgtggtccttgggtgctctggcca  tcgagcggtagctgtgtgtgtgaagccatgagcaactccgcttcgg  ggagaacctgccaatcagggcgttgccttcacctgggtcaggtcggcgtg  gcctgcccgcacccccactcggcggtggtccaggfacatccccga  gggctcagctgctcgtgtggaalcgactactacacgctcaagccgga  ggtaacaacgagcttttctcatctacatgtctgtggtccactcaccatc  cccatgattatcatcttttctctatgggcagctcgtctcaccgtcaagg  aggccgctgccagcagcaggagtcagccaccacacagaaggcag  agaaggaggcaccgccatggtcactcatggtcactcgttctctgatc  tgctgggtgccctacgccagctggcattctacatctcaccaccagg  gctccaactcgggtccatcttcatgacctcccagcgttctttgccaaga  gcccggccatctacaacctgtcatctatcatgatgaacaagcagttc  cggaaactgcatgctcaccaccatctgctgcggcaagaaccactgggt  gacgatgaggcctctgctaccgtgtccaagacggagacgagccaggt  ggccccggcctaa</p>	<i>sapiens</i>
SEQ ID NO: 9 ( <i>cmRHO</i> cDNA)	<p>atgaatggcacagaaggccctaactctacgtgcccttctccaatgcga  cgggtgtggtacgcagcccctcagtagtaccacagtactacctggctg  agccatggcagttctccatgctggccgctacatgttctgctgatcgtg  ctgggcttcccatacaacttctcactgtacgtcaccgtccagcacia  gaagctgcgcacgcctctcaactacatcctgctcaacctagccgtggct  gaccttctcaggtcctaggtggcttaccagcaccctctacacgtcgt  tcacggatattcgtctcgggcccacaggatgcaattggaggcctctt  tgccaccctggggcgtgaaattgccctgtggtccttgggtgctctggcc  atcagcggtagctgtgtgtgtgaagccatgagcaactccgcttcg  gggagaacctgccaatcagggcgttgccttcacctgggtcaggtcgcgt  ggcctgcggcacccccactcggcggtggtccagglacatccccg  agggcctcagctgctcgtgtggaatcactactacacgtcaagccgg  aggtaacaacgagcttctgcatctacatgctgtggtccactcaccat  cccatgattatcatcttttctctatgggcagctcgtctcaccgtcaag  gaggccgctgccagcagcaggagtcagccaccacacagaaggca  gagaaggaggcaccgccatggtcactcatggtcactcgttctctgat</p>	<i>Homo sapiens</i>

Sequence ID Number	Sequence	Species
	<p>ctgctgggtgccctacgccagcgtggcattctacatcttcacccaccag  ggctccaacttcgggccatcttcacatgaccatcccagcgttctttgccaag  agcgccgccatctacaaccctgtcatctatcatgatgaacaagcagtt  ccggaactgcatgctcaccaccatctgctgctggcaagaaccactggg  tgacgatgaggccctctgctaccgtgtccaagacggagacgagccagg  tgccccggcc</p>	
SEQ ID NO: 10 (CBh promoter)	<p>cgttacataacttacggfaaatggcccgcctggctgaccgcccaacga  ccccgcccatgacgtcaatagtaacgccaataggacttccattga  cgtaatgggtggagtatttacggtaaactgccacttggcagtacatca  agtgtatcatatgccaagtacgccccctattgacgtcaatgacggtaaat  ggcccgcctggcattgtgcccagtacatgacctatgggacttctactt  ggcagtacatctacglattagtcacgctatfaccatggcagggtagc  cccacgttetgettcactcctccccatcccccccccccccaacccccaaft  ttgtatttatttttttaattttttgtgcagcgtggggggcggggggggg  ggggggggcgcgccaggcggggggggggggggggcgaaggggcgg  ggcggggcgaggcggagagggtgcggcggcagccaatcagagcgg  cgcgctccgaaaagttctttatggcagggcggcggcggcggcggc  cctataaaaagcgaagcgcgcggcggcgggagtcgctgcgacgct  gccttcgccccgtccccgcctccggccgcctcgcgccgccgcc  cggtctgactgaccgcgttaccacaggtgagcggcgggacgg  cccttctctccgggctgtaattagctgagcaagaggaagggttaag  ggatggttggttggtgggtatfaatgfttaattacctggagcacctgcct  gaaalcaacttttttcaggtt</p>	<i>Gallus gallus</i>
SEQ ID NO: 11 (BGH poly-A)	<p>taagagctcgctgatcagcctcgactgfccttctagtgtccagccatct  gttgtttgccctccccgtgcttcttgacctggaagggtgccactcc  cactgccccttaataaaaatgaggaatgcatcgcattgtctgagtag  gtgtcattctattctgggggtgggtggggcaggacagcaaggggg  aggattgggaagagaatagcaggcatgctgggga</p>	<i>Bos taurus</i>
SEQ ID NO: 12 (U6 promoter)	<p>gagggcctattcccatgattcctcatattgcatatacgatacaaggctg  ttagagagafaattggaatfaattgactgtaaacacaaaagataatgtac  aaaatacgtgacgtagaagtaataattcttgggtagttgcagttttaa  aftatgttttaaatggactatcatatgcttaccgtaactgaaagfattfcg  attcttggctttatatacttgggaaaggacgaaacacc</p>	<i>Homo sapiens</i>

Sequence ID Number	Sequence	Species
SEQ ID NO: 13 (ITR)	<p>cgcgctcgctcgctcactgaggccgccgggcaaaagcccgggctc  ggcgacctttggcgcgccggcctcagtgagcgagcgagcgcgag  agaggagtggccaaactccatcactaggggtcctgt</p>	<i>Adeno-associated virus-2</i>
SEQ ID NO: 14 (sCMV promoter)	<p>actcacggggattccaagtctccaccattgacgtcaatgggagttg  ttttggcaccaaaatcaacgggactttccaaaatgtcgtataaacccgc  cccgtagcgaatggcggtaggcgtgtacgggtgggaggtctat  aagcagagctcgttagtgaaccgt</p>	<i>Human herpesvirus 5</i>
SEQ ID NO: 15 (3xFlag)	<p>GACTATAAGGACCACGACGGAGACTACAA  GGATCATGATATTGATTACAAAGACGATGA  CGATAAG</p>	
SEQ ID NO: 16 (SV40-NLS)	CCAAAGAAGAAGCGGAAGGTC	<i>Simian virus 40</i>
SEQ ID No. 17 (Cas9)	<p>gacaagaagtacagcatcggcctggac  atcggcaccactctgtgggctgggccc  gtgatcaccgacgagfacaagtgccc  agcaagaaatcaaggctciggcgaac  accgaccggcacagcaicaagaaga  cctgatcggagccctgctgtgacagc  ggcgaaacagccgagggccaccggct  gaagagaaccgccagaagaagataca  ccagaccggaagaaccggatctctatct  gcaagagatctcagcaacgagatggc  caagggtgacgacagctctccacaga  ctggaagagctctcctggtggaagg  ataagaagcacgagcggcaccatct  tcggcaacatcgtggacaggtggcct  accacgagaaglacccaccatctacc  acctgagaagaactggtggacagca  ccgacaaggccgacctggcgtgatct  atctggccctggccacatgatcaagttc  cggggccacttctgatcggggcgac  ctgaaccccgacaacagcgagctggac  aagctgttcatccagctggtgcagacct  acaaccagctgtcggagaaaacccat  caaccagcggcgtggacccaagg  ccatcctgtctgccagactgagcaagag  cagacggctggaaaatctgatcgcca  gctgcccggcgagaagaagaatggcct  gttcggaacctgattgccctgagcctg  ggcctgacccccactcaagagcaact  tcgacctggccgaggtgccaactgc  agctgagcaaggacacctcagcagac  gacctggacaacctgtggcccagatc  ggcgaccagctacccgacctgttctgg  ccgcaagaacctgtccgacgccatct</p>	<i>Streptococcus pyogenes</i>

Sequence ID Number	Sequence	Species
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Sequence ID Number	Sequence	Species
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Sequence ID Number	Sequence	Species
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SEQ ID No. 18 (nucleoplasmin NLS)	aaaaggccggcggccacgaaaaaggc cggccaggcaaaaaagaaaaag	<i>Xenopus laevis</i>
SEQ ID No. 19 (synthetic poly A)	aataaaagatcittatitctatagatctgt gtgtgggtttttgtgtg	<i>Oryctolagus cuniculus</i>
SEQ ID No. 20 (CMV promoter)	cgtfacataacttacggtaaatggcccgc ctggctgaccgcccacgaccccgcgc caltgacgtcaataatgacgtatgtccc atagtaacgccaataggactttccattg acgtcaatgggtggagtattacggtaaa ctgccacttggcagttacatcaagtgtat catatgccaaagtacgcccctattgacgt caatgacggtaaatggcccgcctggcat tatgccagttacatgacctatgggacttt cctactggcagttacatctacgtattagtc atcgctattacatgggtgatgcgggtttgg cagtacatcaatggcgtggatagcgggt ttgactcacgggatttccaagtctcac cccattgacgtcaatgggagttgtttgg caccaaaatcaacgggactttcaaaat gtcgtaaacactccgcccattgacgca aatgggcggtagggcgtgtacgggtggga ggtctatataagcagagct	<i>Human herpesvirus 5</i>
SEQ ID No. 21 (EGFP)	atggtagcaaggcggaggagctgtc accggggtgggtcccatectgtgag ctggacggcgacgtaaacggccacaa gttcagcgtgtccggcggaggcggagg gcgatgccacctacggcaagctgacc tgaagttcatctgaccaccggcaagct gccctgtccctggcccaccctcgtgac caccctgacctacggcgtgcafgcttc agccgtaccccgaccacatgaagcag cacgacttctcaagtccgcatgcccg aaggctacgtccaggagcgcaccatctt ctcaaggacgacggcaactacaagac ccgcgccgaggtgaagttcaggggcg acaccctggtgaaccgcatcgagctga agggcatcgacttcaaggaggacggca acatctggggcacaagctggagfaca actacaacagccacaacgtctatatcatg	<i>Aequorea victoria</i>

Sequence ID Number	Sequence	Species
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Sequence ID Number	Sequence	Species
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Sequence ID Number	Sequence	Species
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Sequence ID Number	Sequence	Species
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	<p>TGATGGGCCGGCACAAG  CCCGAGAACATCGTGAT  CGAAATGGCCAGAGAGA  ACCAGACCACCCAGAAG  GGACAGAAGAACAGCCG  CGAGAGAATGAAGCGGA  TCGAAGAGGGCATCAA  GAGCTGGGCAGCCAGAT  CCTGAAAGAACACCCCG  TGGAAAACACCCAGCTG  CAGAACGAGAAGCTGTA  CCTGTACTACCTGCAGA  ATGGGCGGGATATGTAC  GTGGACCAGGAACTGGA  CATCAACCGGCTGTCCG  ACTACGATGTGGACCAT  ATCGTGCCTCAGAGCTTT  CTGAAGGACGACTCCAT  CGACAACAAGGTGCTGA  CCAGAAGCGACAAGAAC  CGGGGCAAGAGCGACAA  CGTGCCCTCCGAAGAGG  TCGTGAAGAAGATGAAG  AACTACTGGCGGCAGCT  GCTGAACGCCAAGCTGA  TTACCCAGAGAAAGTTC  GACAATCTGACCAAGGC  CGAGAGAGGCCGGCCTGA</p>	

Sequence ID Number	Sequence	Species
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	<p>GCGACTACAAGGTGTAC  GACGTGCGGAAGATGAT  CGCCAAGAGCGAGCAGG  AAATCGGCAAGGCTACC  GCCAAGTACTTCTTCTAC  AGCAACATCATGAACTT  TTCAAGACCGAGATTA  CCCTGGCCAACGGCGAG  ATCCGGAAGCGGCCTCT  GATCGAGACAAACGGCG  AAACCGGGGAGATCGTG  TGGGATAAGGGCCGGGA  TTTTGCCACCGTGCGGA  AAGTGCTGAGCATGCC  CAAGTGAATATCGTGAA  AAAGACCGAGGTGCAGA  CAGGCGGCTTCAGCAA  GAGTCTATCCTGCCAA  GAGGAACAGCGATAAGC  TGATCGCCAGAAAGAAG  GACTGGGACCCTAAGAA  GTACGGCGGCTTCGACA  GCCCCACCGTGGCCTAT  TCTGTGCTGGTGGTGGC  CAAAGTGGAAGGGCA  AGTCCAAGAACTGAAG  AGTGTGAAAGAGCTGCT  GGGGATCACCATCATGG  AAAGAAGCAGCTTCGAG  AAGAATCCCATCGACTT  TCTGGAAGCCAAGGGCT  ACAAAGAAGTGA AAAA  GGACCTGATCATCAAGC  TGCCTAAGTACTCCCTGT  TCGAGCTGGAAAACGGC  CGGAAGAGAATGCTGGC  CTCTGCCGGCGAACTGC  AGAAGGGAAACGAACTG</p>	

Sequence ID Number	Sequence	Species
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Sequence ID Number	Sequence	Species
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## SEQUENCE LISTING

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&lt;150&gt; 62/154,966

&lt;151&gt; 2015-04-30

&lt;160&gt; 27

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 aaatcctt 128

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**Patentkrav**

1. Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ved behandling af en autosomt dominant øjensygdom hos et individ, idet anvendelsen omfatter indgivelse til individet af en terapeutisk effektiv mængde af sammensætningen, hvor den mindst ene type af AAV-vektoren omfatter:

(a) to guide-RNA-sekvenser, der hybridiserer med det endogene autosomt dominante sygdomsrelaterede gen hos individet;

(b) en anden sekvens, der omfatter et kodonmodificeret autosomt dominant sygdomsrelateret gen eller fragment deraf, hvor mindst en sygdomsrelateret mutation er rettet i det kodonmodificerede autosomt dominante sygdomsrelaterede gen eller fragment deraf, og hvor det kodonmodificerede autosomt dominante sygdomsrelaterede gen eller fragment deraf ikke genkendes af guide-RNA'et; og

(c) en tredje sekvens, der koder for en Cas-nuklease, eventuelt Cas 9, hvor mindst en guide-RNA-sekvens er valgt fra gruppen bestående af SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7 eller kombinationer deraf.

2. Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor sammensætningen omfatter to typer rekombinante AAV-vektorer, der skal indgives til individet, hvor en første type af rekombinant AAV-vektor omfatter guide-RNA-sekvensen og den anden sekvens, og hvor en anden type af rekombinant AAV-vektor omfatter den tredje sekvens.

3. Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor øjensygdommen er valgt fra gruppen bestående af autosomal dominant korioretinal atrofi eller degeneration, autosomal dominant tap- eller tap-stav-dystrofi, autosomal dominant medfødt stationær

natteblindhed, autosomal dominant Lebers kongenitale amaurose, autosomal dominant makuladegeneration, autosomal dominant okulær-retinal udviklings sygdom, autosomal dominant optisk atrofi, autosomal dominant retinitis pigmentosa, autosomal dominante syndromiske/systemiske sygdomme med retinopati, Sorsbys makuladystrofi,

5 aldersrelateret makuladegeneration, Doynes honeycomb-makulasygdom og juvenil makuladegeneration.

4. Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor øjensygdommen er retinitis pigmentosa, aldersrelateret makuladegeneration eller Doynes honeycomb.

5. Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor det autosomt dominante sygdomsrelaterede gen er RHO-genet eller EFEMP1-genet.

6. Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ifølge krav 1 eller 2, hvor den rekombinante AAV-vektor er en AAV2-vektor eller en AAV8-vektor

7. Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor CRISPR-Cas-systemet styres af en promoter, der styrer ekspressionen af det kodonmodificerede autosomt dominante sygdomsrelaterede gens produkt i øjenceller.

8. Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor (i) det kodonmodificerede autosomt dominante sygdomsrelaterede gen eller fragment deraf er integreret i det

endogene autosomt dominante sygdomsrelaterede gen, eller hvor (ii) det kodonmodificerede autosomt dominante sygdomsrelaterede gen eller fragment deraf ikke er integreret i det endogene autosomt dominante sygdomsrelaterede gen.

- 5 **9.** Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor det autosomt dominante sygdomsrelaterede gen er et gen, der er valgt fra gruppen bestående af PRDM13, RGR, TEAD1, AIPL1, CRX, GUCA1A, GUCY2D, PITPNM3, PROM1, PRPH2, RIMS 1, 10 SEMA4A, UNCI 19, GNAT1, PDE6B, RHO, WSF1, IMPDH1, OTX2, BEST1, C1QTNF5, CTNNA1, EFEMP1, ELOVL4, FSCN2, GUCA1B, HMCN1, IMPG1, RP1L1, TIMP3, VCAN, MFN2, NR2F1, OPA1, ARL3, CA4, HK1, KLHL7, NR2E3, NRL, PRPF3, PRPF4, PRPF6, PRPF8, PRPF31, RDH12, ROM1, RP1, RP9, RPE65, SNRNP200, SPP2, TOPORS, ABCC6, ATXN7, COL11A1, COL2A1, JAG1, KCNJ13, 15 KIF11, OPA3, PAX2, TREX1, CAPN5, CRB 1, FZD4, ITM2B, LRP5, MAPKAPK3, MIR204, OPN1SW, RB I, TSPAN12 og ZNF408.

- 10.** Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant 20 sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor den rekombinante AAV-vektor er egnet til at indgives ved injektion i øjet.

- 11.** Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant 25 sygdomsrelateret gen, til anvendelse ifølge krav 1, hvor det kodonmodificerede autosomt dominante sygdomsrelaterede gen eller fragment deraf er valgt fra gruppen bestående af SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 23, SEQ ID NO: 24, SEQ ID NO: 25, SEQ ID NO: 26 og SEQ ID NO: 27 eller kombinationer deraf.

- 30 **12.** Sammensætning, der omfatter mindst en type af rekombinant adenoassocieret viral (AAV) vektor, som koder for et CRISPR-Cas-system rettet mod et autosomt dominant sygdomsrelateret gen til anvendelse ifølge krav 1, hvor det endogene autosomt

dominante sygdomsrelaterede gen er vildtype og/eller mutant.

DRAWINGS

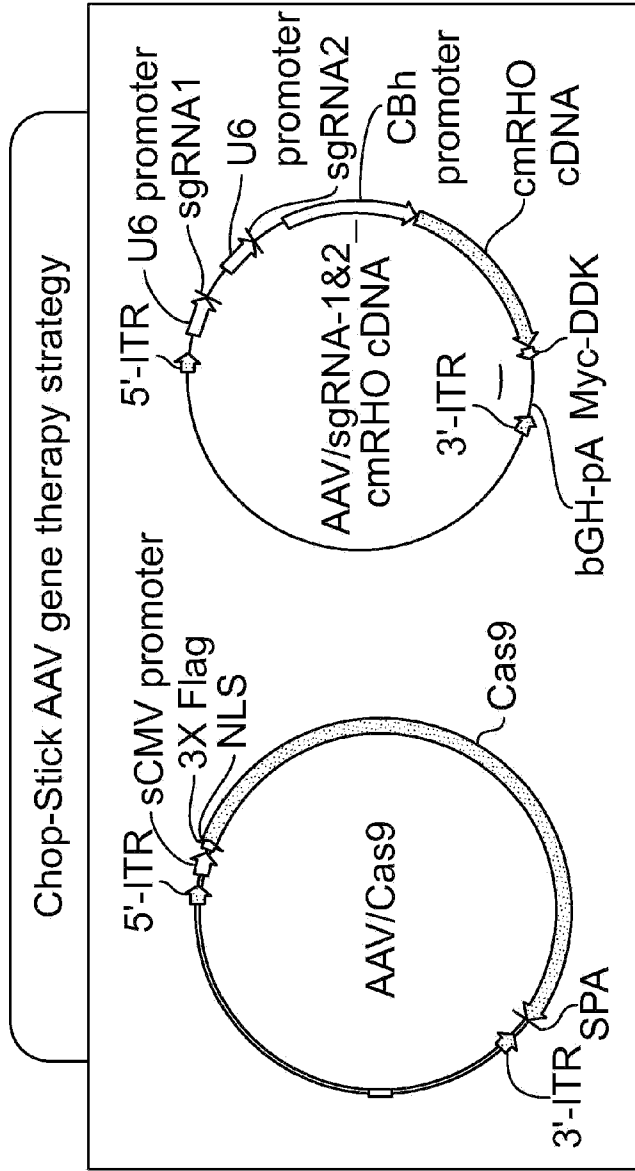


FIG. 1A

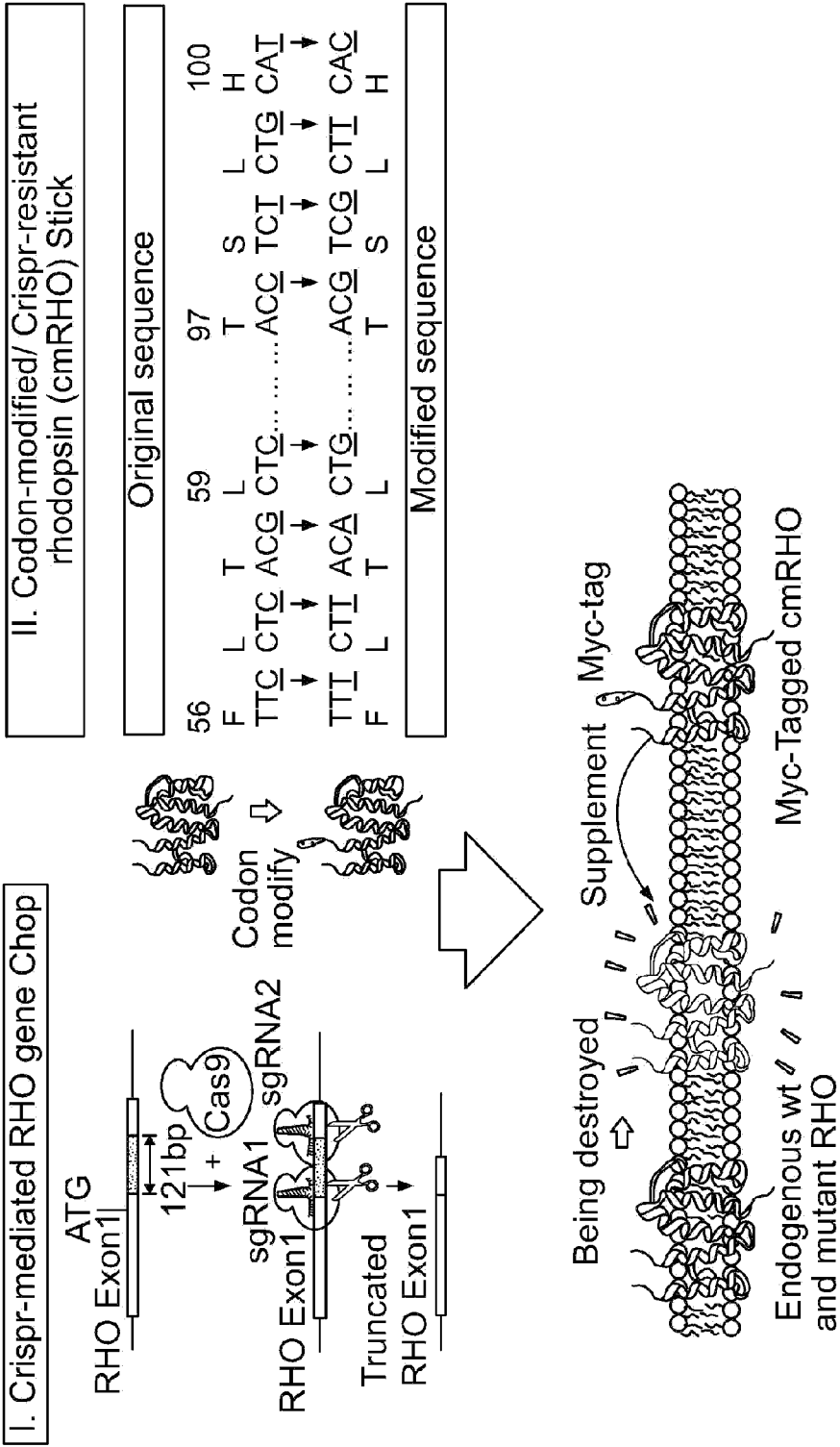


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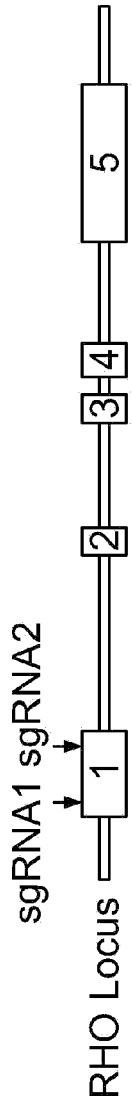


FIG. 2A



FIG. 2B

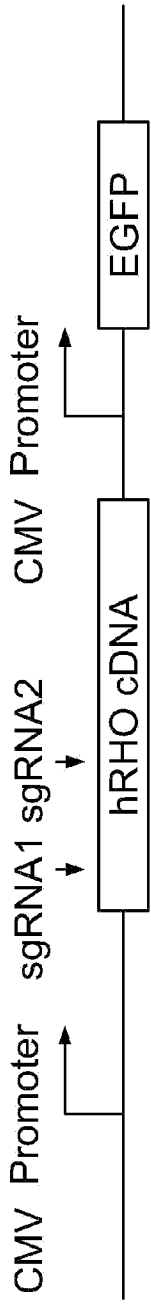


FIG. 3A

Normalized Expression of RHO

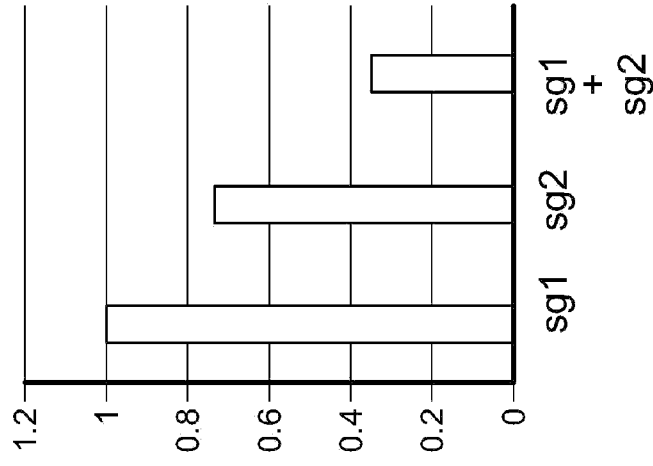


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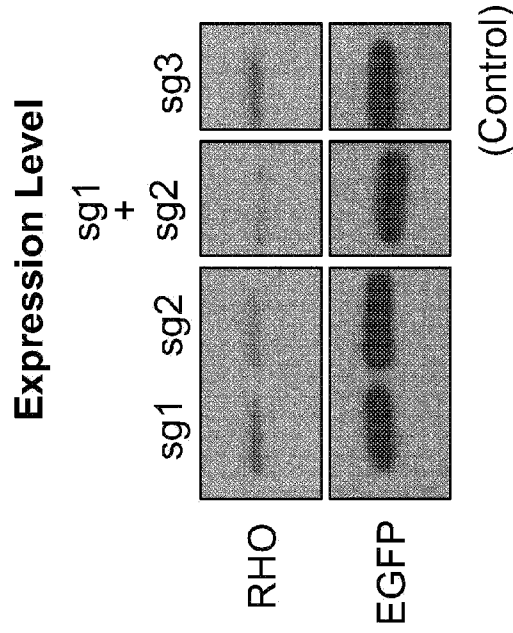


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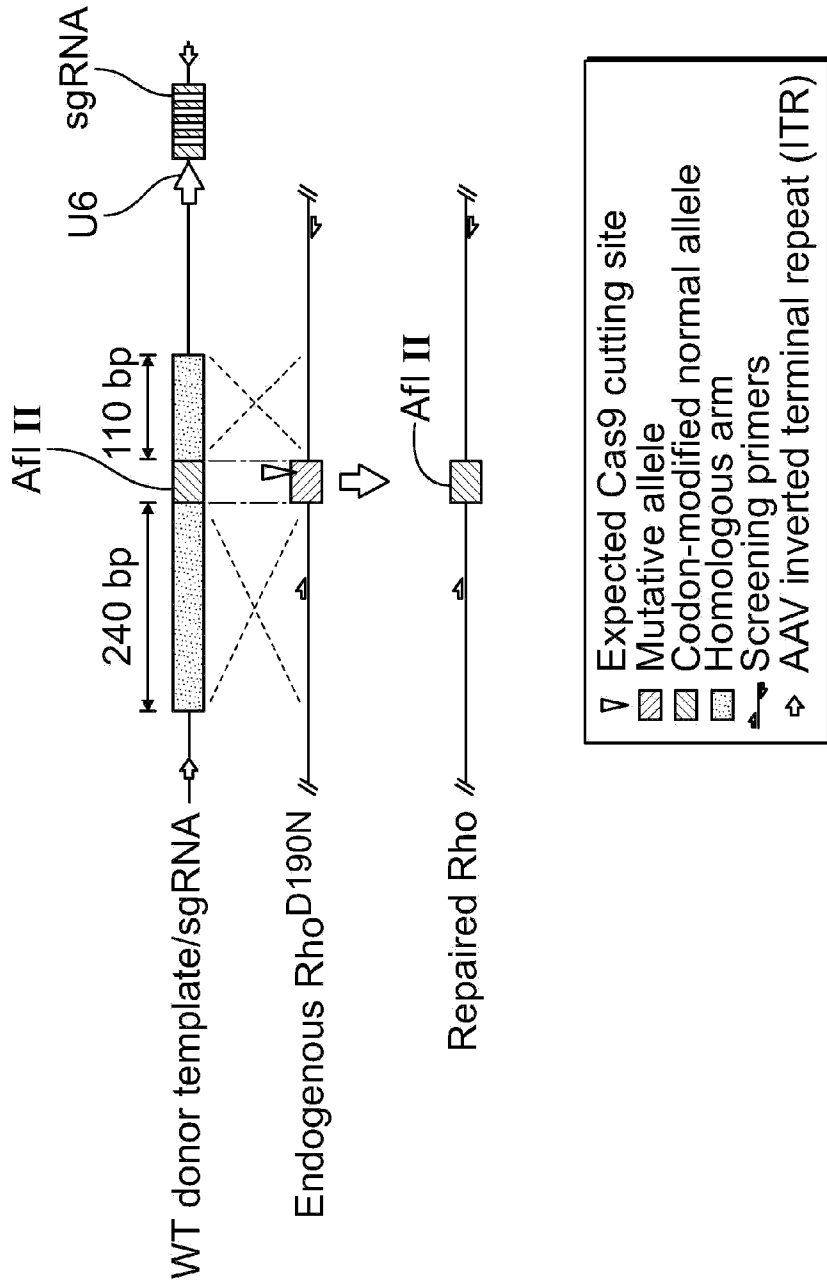


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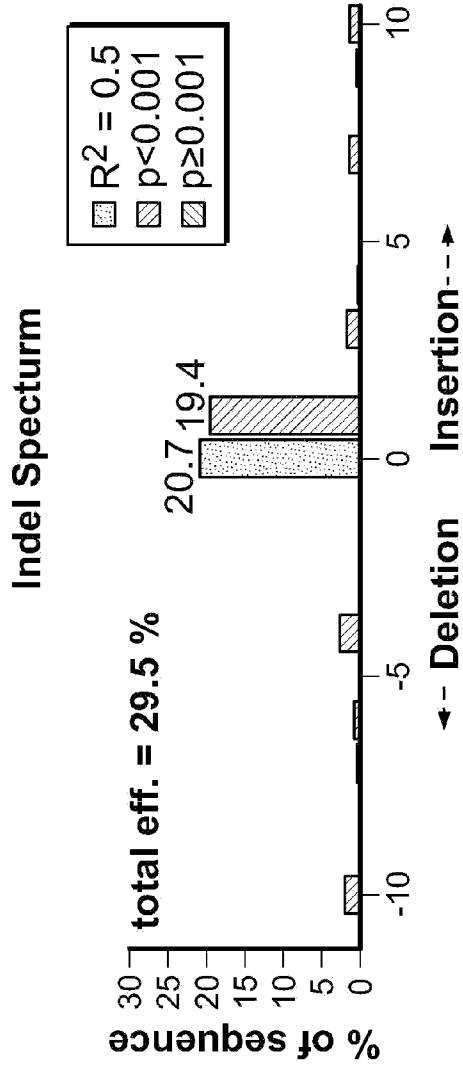


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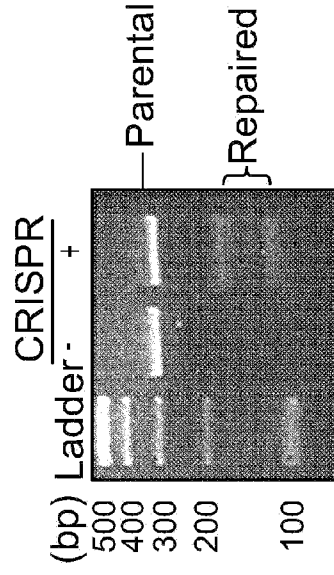


FIG. 4C

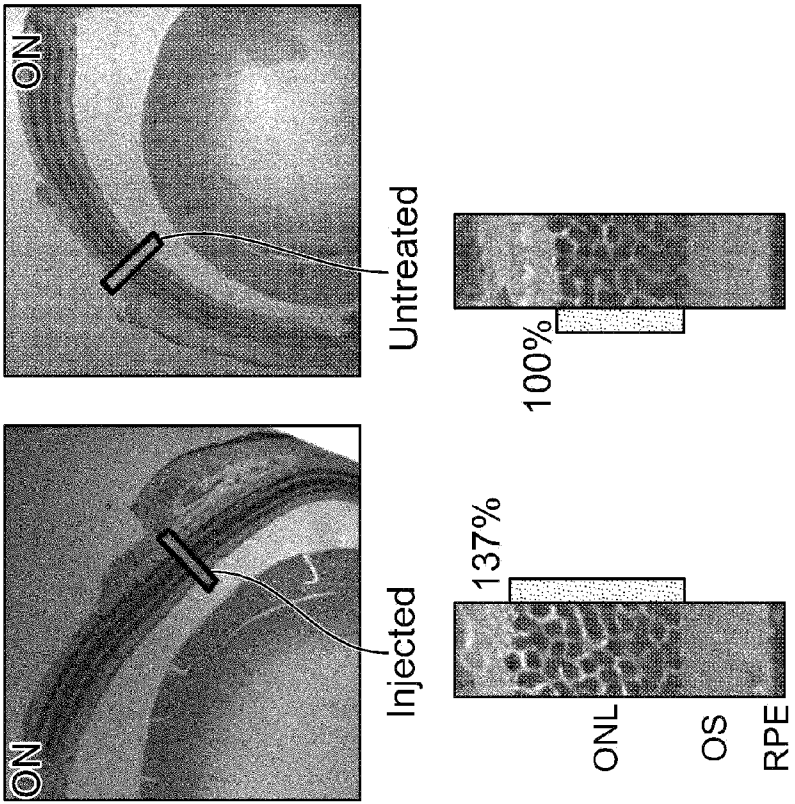


FIG. 5A

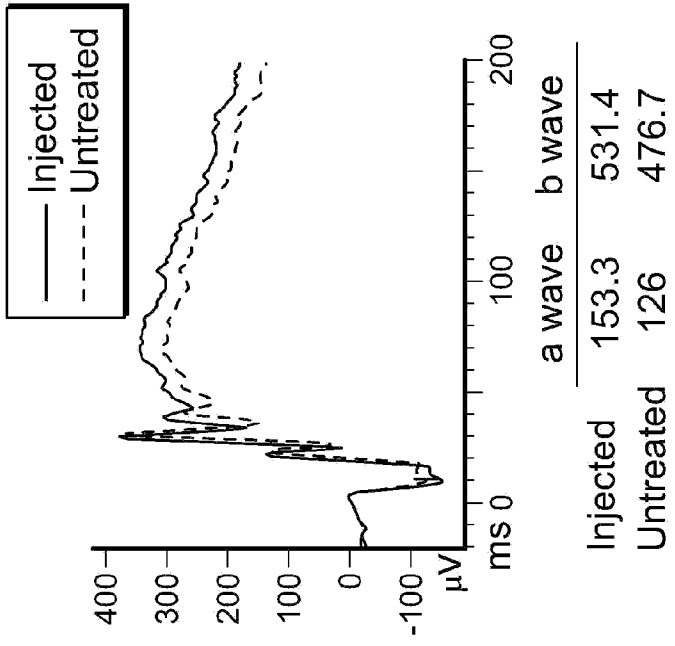


FIG. 5B

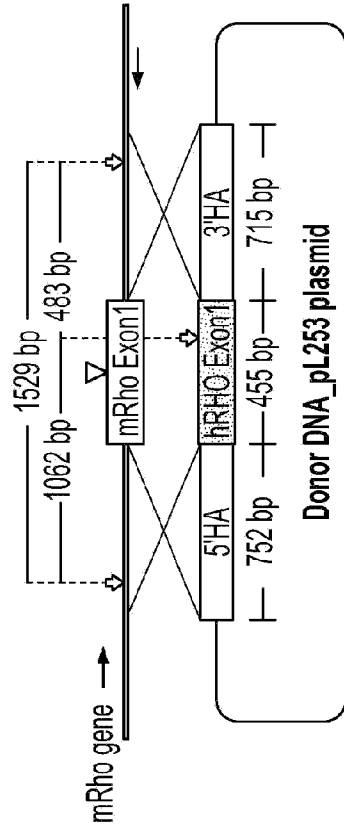


FIG. 6A

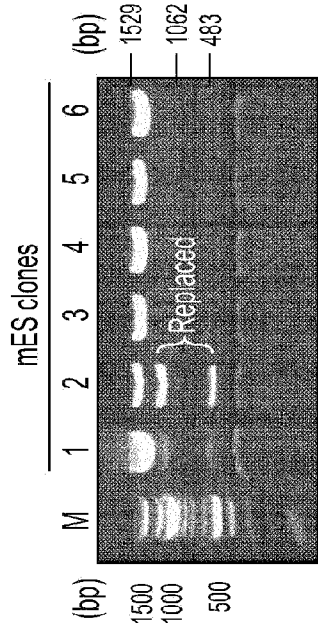


FIG. 6B

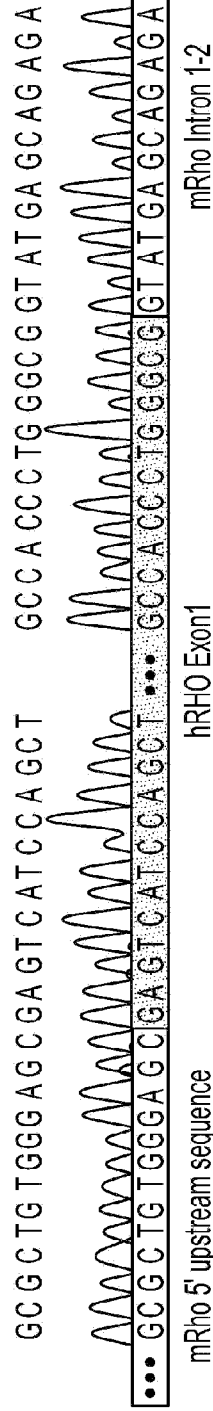


FIG. 6C

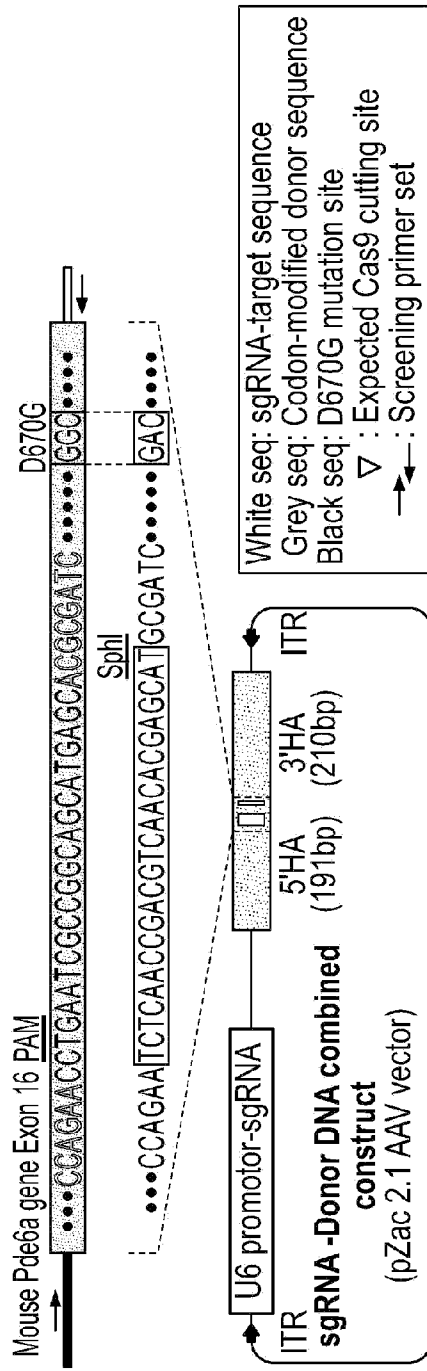


FIG. 7A

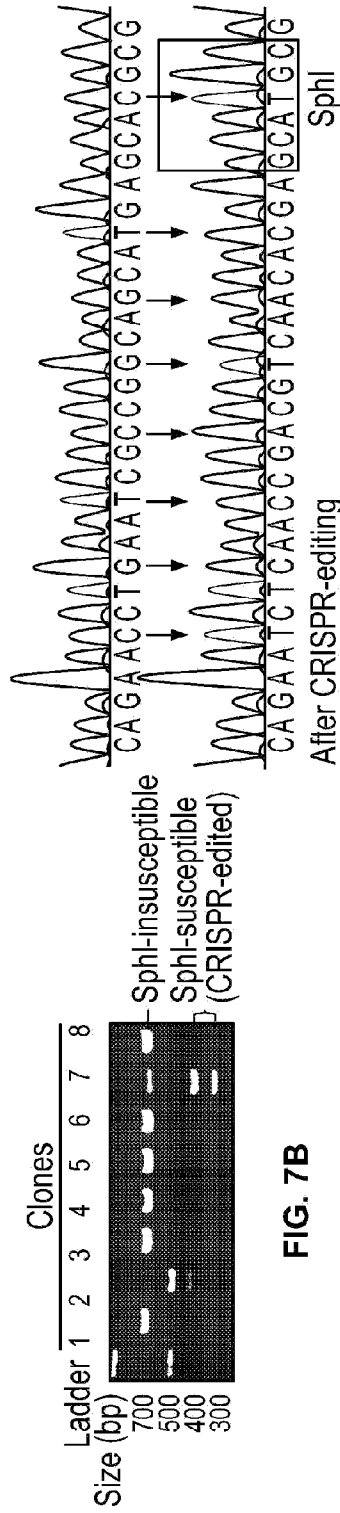


FIG. 7B

FIG. 7C

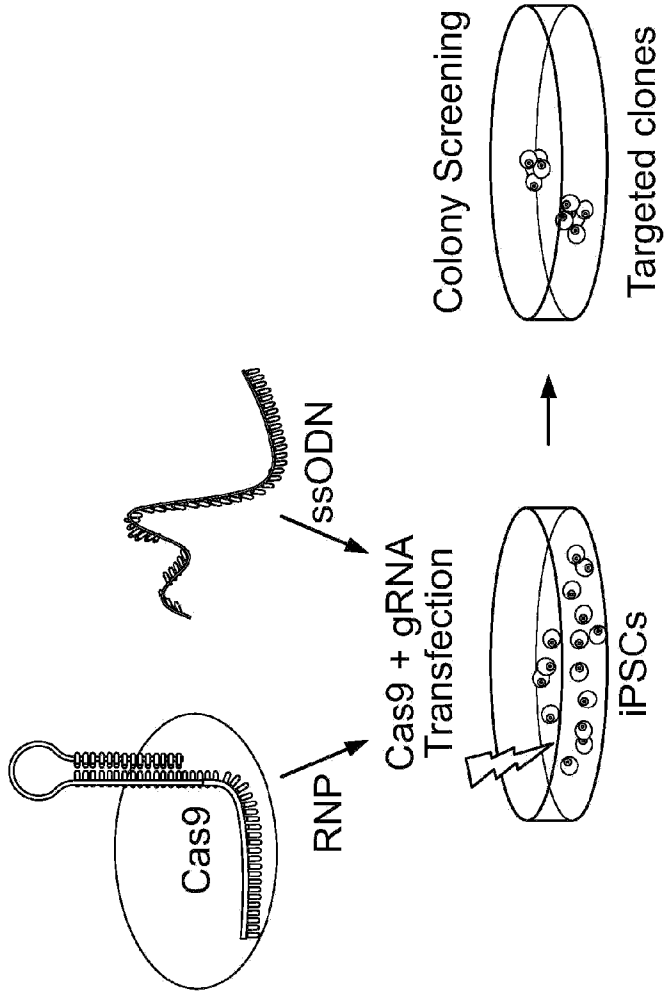


FIG. 8A

gRNA sequence tgagaccacaaatgaatgct

EFEMP1 R345W sequence

cagatataaatgagtgtagaccacaaatgaatgctgggaggatgaaatgtgttggaattatcatggcgg  
 gtctataattactcacactctgggttacttacgaccctcctactttacacaaaccttaatagtagtaccgcc

FIG. 8B



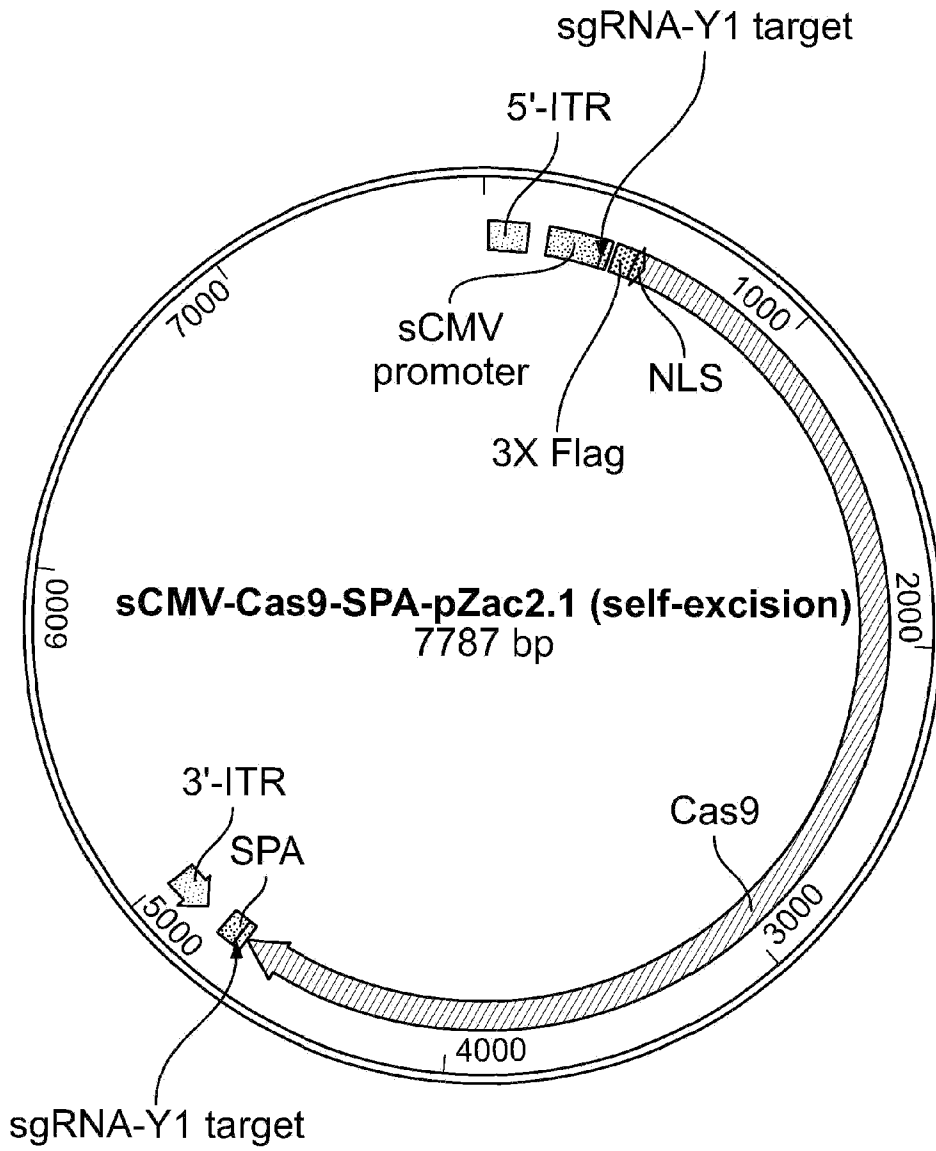


FIG. 9