METHODS AND COMPOSITIONS FOR TREATING BRAIN DISEASES

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CPC .......................... A61K 48/0075 (2013.01); C12N 9/485 (2013.01); C12N 15/86 (2013.01); C12N 2799/025 (2013.01)

Field of Classification Search

CPC .............................. C12N 15/1137; C12N 2310/14 See application file for complete search history.

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(Continued)

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ABSTRACT

The present disclosure provides methods of treating a disease in a non-rodent mammal comprising administering to the cerebrospinal fluid (CSF) of the non-rodent mammal an rAAV2 particle containing a vector comprising a nucleic acid encoding a therapeutic protein inserted between a pair of AAV inverted terminal repeats in a manner effective to infect an ependymal cell in the non-rodent mammal, wherein the ependymal cell secretes the therapeutic protein so as to treat the disease.

24 Claims, 15 Drawing Sheets
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* cited by examiner
AAV eGFP in nonhuman primate brain

Figure 2

4th ventricle

3rd ventricle
ClustalW (v1.83) multiple sequence alignment

2 Sequences Aligned  
Alignment Score = nan
Gaps Inserted = 7  
Conserved Identities = 446

Pairwise Alignment Mode: Slow
Pairwise Alignment Parameters:
  Open Gap Penalty = 10.0  
  Extend Gap Penalty = 0.1
  Similarity Matrix: gonnet

Multiple Alignment Parameters:
  Open Gap Penalty = 10.0  
  Extend Gap Penalty = 0.1
  Delay Divergent = 40%  
  Gap Distance = 8
  Similarity Matrix: gonnet

Processing time: 0.2 seconds

AAV4capPro 1 -MTDGYLPDWWLEDNLSEQVREWALQPGAPKPKANQHQDNARGLVLPGYKYLGPNGLD 59
  **********.*****.* ** * ** .**.**. ********** ****
AAV2capPro 1 MAAGYLPDWWLEDTLSEQGIRQWKLPGPPPPKPAERHKODSRLGLVLPGYKYLGPFGNLD 60
AAV4capPro 60 KGEPVNAADAAALHEHKAYDQQLKGDNFYLYKHADAEMFQQLQGDSGNGNLGRAVQF 119
AAV2capPro 61 KGEPVNEADAALHEDKAYRQLDSGDNFYLYKHADAEMFQERLKDTSGNGNLGRAVFQ 120
  ***** ************.* ** **************.**.************
AAV4capPro 120 AKKRVLEPLGLVEQAGETPGKRRPLIESPQPPDSSTGIGGGKQPAKKLVFEDETGAG 179
AAV2capPro 121 AKKRVLEPLGLVEEPVKTPAGKKRPVEHPVEPDSSTGIGAGQPPARKRLNFGQGTGDAD 180
ClustalW (v1.83) multiple sequence alignment

2 Sequences Aligned  Alignment Score = nan
Gaps Inserted = 10  Conserved Identities = 1440

Pairwise Alignment Mode: Slow
Pairwise Alignment Parameters:
  Open Gap Penalty = 10.0  Extend Gap Penalty = 5.0

Multiple Alignment Parameters:
  Open Gap Penalty = 30.0  Extend Gap Penalty = 5.0
  Delay Divergent = 40%  Transitions: Weighted

Processing time: 1.8 seconds

1. AAV2capNuc vs. AAV4capNucl

  Aligned Length = 2235  Gaps = 10
  Identities = 1440 (65%)

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<tr>
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<th>AAV2capNuc</th>
<th>AAV4capNucl</th>
</tr>
</thead>
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<tr>
<td>1</td>
<td>ATGGCTGCCGATGGTTATCTTCCAGATTGGCTCGAGGACACTCTCTCTGAAGGAATAAGA</td>
<td>---ATGACCTGACGGTTACCTCCAGATTGGCTAGAGGACAACCTCTCTGAAGGCGTTCTGA</td>
</tr>
<tr>
<td></td>
<td>** ****** ******* ******* ******* ******* ******* ****** **</td>
<td></td>
</tr>
<tr>
<td>61</td>
<td>CAGTGGTGGAAGCTCAAAACCCTGACCACCCACCACAAAGCCGAGGCGGCAATAAGGAC</td>
<td>GAGTGGTGCTGGGCTCAACCCTGGAGCCCCCTTAACCCCAAGGCAAATCAACACATCAAGGAC</td>
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<td></td>
<td>******* *** ******* * ** ******* *** ******* * ** ******* *****</td>
<td></td>
</tr>
<tr>
<td>58</td>
<td>GAGTGGTGGAAGCTCAACCTGACCACCCACCACAAAGCCGAGGCGGCAATAAGGAC</td>
<td>******* *** ******* * ** ******* *** ******* * ** ******* *****</td>
</tr>
</tbody>
</table>

Figure 6B-1
Figure 6B-2

AAV2capNuc 121 GACAGCAGGGTCTTTGTGCTTTCTGCTGAACGACCTGCACTCGAC 180
AAV4capNuc 118 AACGTCGAGTCGCTTTCTACTTCAACCGACGACTCGAC 177

** *********** ** **** *********** ***********

AAV2capNuc 181 AAGGGAGAGCCGTCACGACGGACACCGAGCGGAGCAGGTCTACGAC 240
AAV4capNuc 178 AAGGGGCAACCCCGTCACGGCAGCGGAGCAGCGGAGCAGGTCTACGAC 237

***** ** ** ******** ** **** ** *********** ***********

AAV2capNuc 241 CGGCAGCTCAGACCGGAGAACAACCGTACCTCAAGTACAACACCGGAGCGGAGTTT 300
AAV4capNuc 238 CGGCAGCTCAGACCGGAGAACAACCGTACCTCAAGTACAACACCGGAGCGGAGTTT 297

* ******** * ******** *** ********** ***********

AAV2capNuc 301 CAGGAGCGCTAAAGAGATACGCTTTTGGGCAACCTCGAGACGACTCTCCAG 360
AAV4capNuc 298 CAGGAGCGCTAAAGAGATACGCTTTTGGGCAACCTCGAGACGACTCTCCAG 357

***** *** *** * *** ** *********** ***********

AAV2capNuc 361 GCCAAAAAGAGGGTTTCTGAAACTCTCGGCTTGAGGAACTGTGTAAAGACGCCCTCG 420
AAV4capNuc 358 GCCAAAAAGAGGGTTTCTGAAACTCTCGGCTTGAGGAACTGTGTAAAGACGCCCTCG 417

** *********** ** ** ** ********** ***********

AAV2capNuc 421 GAAAAAAAGAGCGCGTTAGACAGACTCCTCGGAGGACAGACTCTCCCTCGGGAACCGGA 480
AAV4capNuc 418 GAAAAAAAGAGCGCGTTAGACAGACTCCTCGGAGGACAGACTCTCCCTCGGGAACCGGA 477

***** **** ** ** ** ** *********** ** **

AAV2capNuc 481 AAGGCGGGCCAGACGCTGGCAAGAAAAAGATTGGATTGTTTTGTCAGACTGGAGACGCAGAC 540
AAV4capNuc 478 AAAAAAGGCAAGCGGCCGCTAAAAGAAAAGACGCTTTTCTGA---------AGACGAAACT 528

** *** ******** ** ** ** * ** **

***** *
Figure 6B-5

AAV2capNuc  1372  TCAAA--GGCTTCAGTTTTTCAGCCAGCAGGC--AGTGACATTTCGGGACCAGGTCTAGGAAC  1428
AAV4capNuc  1354  ACTGCCAACCACCAACTTTACCAAGCTGGCCCTACCAACTTTCCACTTTAATAAAAAGAAC  1413
*  *  *  ****  **  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *

AAV2capNuc  1429  TGGGCTTTCCTGGACCCTTACCGCCAGCAGCGAGTATCAAAAGACATCTGGGATAACAAC  1488
AAV4capNuc  1414  TGGGCTTGGCCGGGCTTCAATCAAGCAGCAGGCTTCTCAAGACTGCCAATCAAAACTAC  1473
**  *  ***  *  ***  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *

AAV2capNuc  1489  AACAG---TGAAATCTCCTGGGACTGGGACTACACAGGTACCAAGGTACCA---------CCTCAAT  1533
AAV4capNuc  1474  AAGATCCCCTCAGCCCGGCTGAGCAACTTCAATGCAGACACGACACAGCAGCTCTGGAGC  1533
**  *  ***  *  **  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *

AAV2capNuc  1534  GGCAGAGACTCTCTGTGAAATCCGGCAGCCCGCCACCAAGGAGACCTGAGAA  1593
AAV4capNuc  1534  GGAAGATGGAAGGCTGGCTGACCCTCCTCAACCAGGCCGCATGCGTACCTGGAGAGC  1593
**  *  ***  *  ***  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *

AAV2capNuc  1594  AAGTTTTTTCTCAGAGCGGGTTTCTCATCCTTGGGAAGCAAGGCTGACAGAAAACAAAT  1653
AAV4capNuc  1594  AAGTTTCAG---CAACAGCCAGCTCTCCCTTGGGCCCCTAAACAGCAAGCAGCCACCGGC  1650
*****  *  ***  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *

AAV2capNuc  1654  GTGGGACATTGAAAGAGTGTCTAGTTAGGACAGAAGGAAATGCAAGAACACCAACTCCCCGTTG  1713
AAV4capNuc  1651  ACCGTACCCGGGACTCTGTATCTTACCTCTGAGGAGGAGCTGCGAGCCACCACGCAACC  1710
*  *  *  ***  *  ***  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *

AAV2capNuc  1714  GCTACCGGAGGATATGTTTCTGTATCTACCAACCTCCAGAGGGAACACGCAACACACAGCCT  1773
AAV4capNuc  1711  GATACTGAGGATCTGCGGGCAACCTACCTCGGGGCTGACAGCAGCAAACAGCAACACTGCGGACC  1770
*  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *  *
METHODS AND COMPOSITIONS FOR TREATING BRAIN DISEASES

PRIORITY OF INVENTION

This application claims priority to U.S. Provisional Application No. 61/470,460 that was filed on Mar. 31, 2011. The entire content of this provisional application is hereby incorporated herein by reference.

FEDERAL GRANT SUPPORT

This invention was made with government support under NS068099 awarded by National Institutes of Health. The government has certain rights in the invention.

BACKGROUND

Gene transfer is now widely recognized as a powerful tool for analysis of biological events and disease processes at both the cellular and molecular level. More recently, the application of gene therapy for the treatment of human diseases, either inherited (e.g., ADA deficiency) or acquired (e.g., cancer or infectious disease), has received considerable attention. With the advent of improved gene transfer techniques and the identification of an ever expanding library of defective gene-related diseases, gene therapy has rapidly evolved from a treatment theory to a practical reality.

Traditionally, gene therapy has been defined as a procedure in which an exogenous gene is introduced into the cells of a patient in order to correct an inborn genetic error. Although more than 4500 human diseases are currently classified as genetic, specific mutations in the human genome have been identified for relatively few of these diseases. Until recently, these rare genetic diseases represented the exclusive targets of gene therapy efforts. Accordingly, most of the NIH approved gene therapy protocols to date have been directed toward the introduction of a functional copy of a defective gene into the somatic cells of an individual having a known inborn genetic error. Only recently, have researchers and clinicians begun to appreciate that most human cancers, certain forms of cardiovascular disease, and many degenerative diseases also have important genetic components, and for the purposes of designing novel gene therapies, should be considered “genetic disorders.” Therefore, gene therapy has more recently been broadly defined as the correction of a disease phenotype through the introduction of new genetic information into the affected organism.

In vivo gene therapy, a transferred gene is introduced into cells of the recipient organism in situ that is, within the recipient. In vivo gene therapy has been examined in several animal models. Several recent publications have reported the feasibility of direct gene transfer in situ into organs and tissues such as muscle, hematopoietic stem cells, the arterial wall, the nervous system, and lung. Direct injection of DNA into skeletal muscle, heart muscle and injection of DNA-lipid complexes into the vasculature also has been reported to yield a detectable expression level of the inserted gene product(s) in vivo.

Treatment of diseases of the central nervous system, e.g., inherited genetic diseases of the brain, remains an intractable problem. Examples of such are the lysosomal storage diseases. Collectively, the incidence of lysosomal storage diseases (LSD) is 1 in 10,000 births world wide, and in 65% of cases, there is significant central nervous system (CNS) involvement. Proteins deficient in these disorders, when delivered intravenously, do not cross the blood-brain barrier, or, when delivered directly to the brain, are not widely distributed. Thus, therapies for the CNS deficits need to be developed.

SUMMARY

The present invention provides a method of delivering a nucleic acid to an ependymal cell of a non-rodent mammal comprising administering to the ependymal cell an AAV2 particle containing a vector comprising the nucleic acid inserted between a pair of AAV2 inverted terminal repeats, thereby delivering the nucleic acid to the ependymal cell. In certain embodiments, the rAAV2 particle infects the non-primate ependymal cell at an rate of more than 20% than the infectivity rate of AAV4, such as at a rate of more than 50% or 100%, 1000% or 2000% than the infectivity rate of AAV4.

The present invention provides a method of delivering a nucleic acid to a non-rodent mammal comprising administering to an ependymal cell from the mammal an AAV2 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, and returning the ependymal cell to the mammal, thereby delivering the nucleic acid to the mammal.

The present invention provides a method of delivering a nucleic acid to an ependymal cell in a non-rodent mammal comprising administering to the mammal an AAV2 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to an ependymal cell in the mammal.

The present invention provides a method to deliver an agent to the central nervous system of a non-rodent mammal, comprising administering to the cerebrospinal fluid (CSF) of the non-rodent mammal an AAV2 particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats in a manner effective to infect ependymal cells in the non-rodent mammal such that the ependymal cells secrete the agent into the CSF of the non-rodent mammal.

The present invention provides a method of treating a disease in a non-rodent mammal comprising administering to the ependymal cells of the mammal an AAV2 particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to the ependymal cell.

In certain embodiments, the disease is a lysosomal storage disease (LSD). In certain embodiments, the LSD is infantile or late infantile ceroid lipofuscinosis, neuropathic Gaucher, Juvenile Batten, Fabry, MLD, Sanfilippo A, Hunter, Krabbe, Niemann-Pick C, Tay-Sachs, Hurler (MPS-1 H), Sanfilippo B, Maroteaux-Lamy, Niemann-Pick A, Cystinosis, Hurler-Scheie (MPS-I H/S), Sly Syndromes (MPS VII), Scheie (MPS-I S), Infantile Batten, GM1 Gangliosidosis, Mucolipidosis type II/III, or Sandhoff disease. In certain embodiments, the disease is LINCL. In certain embodiments, the disease is a neurodegenerative disease, such as Huntington’s disease, ALS, hereditary spastic hemiplegia, primary lateral sclerosis, spinal muscular atrophy, Kennedy’s disease, Alzheimer’s disease, a polyglutamine repeat disease, or Parkinson’s disease.

In certain embodiments, the large mammal is a primate, horse, sheep, goat, pig, or dog. In certain embodiments, the primate is a human.

In certain embodiments, the nucleic acid is a lysosomal hydrolase. In certain embodiments, the nucleic acid is TPP1.
The present invention provides a method of transfecting an ependymal cell a non-roden mammalian brain comprising administering to the cerebrospinal fluid (CSF) of the non-roden mammal an AAV2 particle containing a vector comprising a nucleic acid inserted between a pair of AAV2 inverted terminal repeats in a manner effective to infect ependymal cells in the non-roden mammal such that the ependymal cells secrete the agent into the CSF of the non-roden mammal.

The present invention provides a use of the viral vector described hereinabove to prepare a medicament useful for treating a lysosomal storage disease in a mammal.

The present invention provides a cell as described hereinabove for use in medical treatment or diagnosis.

The present invention provides a use of the cell as described hereinabove to prepare a medicament useful for treating a lysosomal storage disease in a mammal.

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 shows the transfection of AAVgGFP in dog.
Fig. 2 shows the transfection of AAVGFP in nonhuman primate brain.
Fig. 3 shows ependymal transduction of TPP1 in NHP brain, indicated significant increase of enzyme in the CSF.
Fig. 4 shows elevated TPP1 activity in various brain regions.
Fig. 5 shows the results of T-maze performance of control and treated dogs. Light circles are for affected dogs; dark squares are for normal dogs, and dark circles are for a TPP−/− dog treated with AAV2-CLN2.
Fig. 6A is an alignment of AAV2 (SEQ ID NO:1) and AAV4 (SEQ ID NO:2) proteins and Fig. 6B is and alignment of AAV2 (SEQ ID NO:3) and AAV4 (SEQ ID NO:4) nucleotides based on the sequence from AAV2 (NC_001401) and AAV4 (NC_001829).

DETAILED DESCRIPTION

Adeno associated virus (AAV) is a small non-pathogenic virus of the parvoviridae family. AAV is distinct from the other members of this family by its dependence upon a helper virus for replication. In the absence of a helper virus, AAV may integrate in a locus specific manner into the q arm of chromosome 19. The approximately 5 kb genome of AAV consists of one segment of single stranded DNA of either plus or minus polarity. The ends of the genome are short inverted terminal repeats which can fold into hairpin structures and serve as the origin of viral DNA replication. Physically, the parvovirus is non-enveloped and its icosahedral capsid is approximately 20 nm in diameter.

To-date eight serologically distinct AAVs have been identified and five have been isolated from humans or primates and are referred to as AAV types 1-5. Govindasamy et al., “Structurally Mapping the Diverse Phenotype of Adeno-Associated Virus Serotype 4,” J. Virol. 80 (23):11556-11570 (2006). The genome of AAV2 is 4680 nucleotides in length and contains two open reading frames (ORFs). The left ORF encodes the non-structural Rep proteins, Rep 40, Rep 52, Rep 68 and Rep 78, which are involved in regulation of replication and transcription in addition to the production of single-stranded progeny genomes. Furthermore, two of the Rep proteins have been associated with the preferential integration of AAV genomes into a region of the q arm of human chromosome 19. Rep68/78 has also been shown to possess NTP binding activity as well as DNA and RNA helicase activities. The Rep proteins possess a nuclear localization signal as well as several potential phosphorylation sites. Mutation of one of these kinase sites resulted in a loss of replication activity.

The ends of the genome are short inverted terminal repeats (ITR) which have the potential to fold into T-shaped hairpin structures that serve as the origin of viral DNA replication. Within the ITR region two elements have been described which are central to the function of the ITR, a GAGC repeat motif and the terminal resolution site (trs). The repeat motif has been shown to bind Rep when the ITR is in either a linear or hairpin conformation. This binding serves to position Rep68/78 for cleavage at the trs which occurs in a site- and strand-specific manner. In addition to their role in replication, these two elements appear to be central to viral integration. Contained within the chromosome 19 integration locus is a Rep binding site with an adjacent trs. These elements have been shown to be functional and necessary for locus specific integration.

The AAV2 virion is a non-enveloped, icosahedral particle approximately 25 nm in diameter, consisting of three related proteins referred to as VP1, VP2 and VP3. The right ORF encodes the capsid proteins VP1, VP2, and VP3. These proteins are found in a ratio of 1:1:10 respectively and are all derived from the right-hand ORF. The capsid proteins differ from each other by the use of alternative splicing and an unusual start codon. Deletion analysis has shown that removal or alteration of VP1 which is translated from an alternatively spliced message results in a reduced yield of infectious particles. Mutations within the VP3 coding region result in the failure to produce any single-stranded progeny DNA or infectious particles. An AAV2 particle is a viral particle comprising an AAV2 capsid protein. An AAV2 capsid polypeptide can encode the entire VP1, VP2 and VP3 polypeptide. The particle can be a particle comprising AAV2 and other AAV capsid proteins (i.e., a chimeric protein, such as AAV4 and AAV2). Variations in the amino acid sequence of the AAV2 capsid protein are contemplated herein, as long as the resulting viral particle comprises the AAV2 capsid remains antigenically or immunologically distinct from AAV4, as can be routinely determined by standard methods. Specifically, for example, ELISA and Western blots can be used to determine whether a viral particle is antigenically or immunologically distinct from AAV4. Furthermore, the AAV2 viral particle preferably retains tissue tropism distinct from AAV4.

An AAV2 particle is a viral particle comprising an AAV2 capsid protein. An AAV2 capsid polypeptide encoding the entire VP1, VP2, and VP3 polypeptide can overall have at least about 63% homology (or identity) to the polypeptide having the amino acid sequence encoded by nucleotides set forth in SEQ ID NO:1 (AAV2 capsid protein). The capsid protein can have about 70% homology, about 75% homology, 80% homology, 85% homology, 90% homology, 95% homology, 98% homology, 99% homology, or even 100% homology to the protein set forth in SEQ ID NO:1. The particle can be a particle comprising both AAV4 and AAV2 capsid protein, i.e., a chimeric protein. Variations in the amino acid sequence of the AAV2 capsid protein are contemplated herein, as long as the resulting viral particle comprising the AAV2 capsid remains antigenically or immunologically distinct from AAV4, as can be routinely determined by standard methods. Specifically, for example, ELISA and Western blots can be used to determine whether a viral
particle is antigenically or immunologically distinct from AAV-4. Furthermore, the AAV2 viral particle preferably retains tissue tropism distinction from AAV-4, such as that exemplified in the examples herein, though an AAV2 chimeric particle comprising at least one AAV2 coat protein may have a different tissue tropism from that of an AAV2 particle consisting only of AAV2 coat proteins.

As indicated in FIGS. 6A and 6B, AAV2 capsid sequence and AAV4 capsid sequence are about 60% homologous. In certain embodiments, the AAV2 capsid comprises (or consists of) a sequence that is at least 65% homologous to the amino acid sequence set forth in SEQ ID NO:1.

In certain embodiments, the invention further provides an AAV2 particle containing, i.e., encapsidating, a vector comprising a pair of AAV2 inverted terminal repeats. The nucleotide sequence of AAV2 ITRs is known in the art. Furthermore, the particle can be a particle comprising both AAV4 and AAV2 capsid protein, i.e., a chimeric protein. Moreover, the particle can be a particle encapsidating a vector comprising a pair of AAV inverted terminal repeats from other AAVs (e.g., AAV1-AAV8). The vector encapsidated in the particle can further comprise an exogenous nucleic acid inserted between the inverted terminal repeats.

The features of AAV have made it an attractive vector for gene transfer. AAV vectors have been shown in vitro to stably integrate into the cellular genome; 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. Integration of AAV provirus is not associated with any long term negative effects on cell growth or differentiation. The ITRs have been shown to be the only cis elements required for replication, packaging and integration and may contain some promoter activities.

The present invention provides methods of administering AAV2 particles, recombinant AAV2 vectors, and recombinant AAV2 virions. An AAV2 particle is a viral particle comprising an AAV2 capsid protein. A recombinant AAV2 vector is a nucleic acid construct that comprises at least one unique nucleic acid of AAV2. A recombinant AAV2 virion is a particle containing a recombinant AAV2 vector. To be considered within the term “AAV2 ITRs” the nucleotide sequence must contain one or both features described herein that distinguish the AAV2 ITR from the AAV4 ITR: (1) three (rather than four as in AAV4) “GAGC” repeats and (2) in the AAV2 ITR Rep binding site the fourth nucleotide in the first two “GAGC” repeats is a C rather than a T.

The promoter can be any desired promoter, selected by known considerations, such as the level of expression of a nucleic acid functionally linked to the promoter and the cell type in which the vector is to be used. Promoters can be an exogenous or an endogenous promoter. Promoters can include, for example, known strong promoters such as SV40 or the inducible metallothionein promoter, or an AAV promoter, such as an AAV p5 promoter. Additional examples of promoters include promoters derived from actin genes, immunoglobin genes, cytomegalovirus (CMV), adenovirus, bovine papilloma virus, adenoviral promoters, such as the adenoviral major late promoter, an inducible heat shock promoter, respiratory syncytial virus, Rous sarcoma virus (RSV), etc. Specifically, the promoter can be AAV2 p5 promoter or AAV4 p5 promoter. Furthermore, smaller fragments of p5 promoter that retain promoter activity can readily be determined by standard procedures including, for example, constructing a series of deletions in the p5 promoter, linking the deletion to a reporter gene, and determining whether the reporter gene is expressed, i.e., transcribed and/or translated.

The AAV2 vector can further comprise an exogenous (heterologous) nucleic acid functionally linked to the promoter. By “heterologous nucleic acid” is meant that any heterologous or exogenous nucleic acid can be inserted into the vector for transfer into a cell, tissue or organism. The nucleic acid can encode a polypeptide or protein or an antisense RNA, for example. By “functionally linked” is meant such that the promoter can promote expression of the heterologous nucleic acid, as is known in the art, such as appropriate orientation of the promoter relative to the heterologous nucleic acid. Furthermore, the heterologous nucleic acid preferably has all appropriate sequences for expression of the nucleic acid, as known in the art, to functionally encode, i.e., allow the nucleic acid to be expressed. The nucleic acid can include, for example, expression control sequences, such as an enhancer, and necessary information processing sites, such as ribosome binding sites, RNA splicing sites, polyadenylation sites, and transcriptional terminator sequences.

The heterologous nucleic acid can encode beneficial proteins that replace missing or defective proteins required by the subject into which the vector is transferred or can encode a cytotoxic polypeptide that can be directed, e.g., to cancer cells or other cells whose death would be beneficial to the subject. The heterologous nucleic acid can also encode antisense RNAs that can bind to, and thereby inactivate, mRNAs made by the subject that encode harmful proteins. In one embodiment, antisense polynucleotides can be produced from a heterologous expression cassette in an AAV2 viral construct where the expression cassette contains a sequence that promotes cell-type specific expression.

Examples of heterologous nucleic acids which can be administered to a cell or subject as part of the present AAV2 vector can include, but are not limited to the nucleic acids encoding therapeutic agents, such as lysosomal hydrolases; tumor necrosis factors (TNF), such as TNF-alpha; interferons, such as interferon-alpha, interferon-beta, and interferon-gamma; interleukins, such as IL-1, IL-1beta, and IL-2 through -14; GM-CSF; adenosine deaminase; secreted factors such as growth factors; ion channels; chemotherapeutics; lysosomal proteins; anti-apoptotic gene products; proteins promoting neural survival such as glutamate receptors and growth factors; cellular growth factors, such as lymphokines; soluble CD4; Factor VIII; Factor IX; T-cell receptors; LDL receptor; ApoE; ApoC; alpha-l antitrypsin; ornithine transcarbamylase (OTC); cystic fibrosis transmembrane receptor (CFTR); insulin; Fc receptors for antigen binding domains of antibodies, such as immunoglobulins; and antisense sequences which inhibit viral replication, such as antisense sequences which inhibit replication of hepatitis B or hepatitis non-A, non-B virus. Furthermore, the nucleic acid can encode more than one gene product, limited only by the size of nucleic acid that can be packaged.

An AAV2 particle is a viral particle comprising an AAV2 capsid protein. Variations in the amino acid sequence of the AAV2 capsid protein are contemplated herein, as long as the resulting viral particle comprising the AAV2 capsid remains antigenically or immunologically distinct from AAV4, as can be routinely determined by standard methods. Specifically, for example, ELISA and Western blots can be used to determine whether a viral particle is antigenically or immunologically distinct from other AAV serotypes.

The term “polypeptide” as used herein refers to a polymer of amino acids and includes full-length proteins and frag-
ments thereof. Thus, "protein," "polypeptide," and "peptide" are often used interchangeably herein. Substitutions can be selected by known parameters to be neutral. As will be appreciated by those skilled in the art, the invention also includes those polypeptides having slight variations in amino acid sequences or other properties. Such variations may arise naturally as allelic variations (e.g., due to genetic polymorphism) or may be produced by human intervention (e.g., by mutagenesis of cloned DNA sequences), such as induced point, deletion, insertion and substitution mutants.

Minor changes in amino acid sequence are generally preferred, such as conservative amino acid replacements, small internal deletions or insertions, and additions or deletions at the ends of the molecules. These modifications can result in changes in the amino acid sequence, provide silent mutations, modify a restriction site, or provide other specific mutations.

The present method provides a method of delivering a nucleic acid to a cell comprising administering to the cell an AAV2 particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to the cell. Administration to the cell can be accomplished by any means, including simply contacting the particle, optionally contained in a desired liquid such as tissue culture medium, or a buffered saline solution, with the cells. The particle can be allowed to remain in contact with the cells for any desired length of time, and typically the particle is administered and allowed to remain indefinitely. For such in vitro methods, the virus can be administered to the cell by standard viral transduction methods, as known in the art and as exemplified herein. Titers of virus to administer can vary, particularly depending upon the cell type, but will be typical of that used for AAV transduction in general. Additionally the titers used to transduce the particular cells in the present examples can be utilized. The cells can include any desired cell in humans as well as other large (non-rodent) mammals, such as primates, horse, sheep, goat, pig, and dog.

More specifically, the present invention provides a method of delivering a nucleic acid to an epidermal cell, comprising administering to the epidermal cell an AAV2 particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to the epidermal cell.

The present invention also includes a method of delivering a nucleic acid to a subject comprising administering to a cell from the subject an AAV2 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, and returning the cell to the subject, thereby delivering the nucleic acid to the subject. The AAV1TRs can be AAV2 ITRs. For such an ex vivo administration, cells are isolated from a subject by standard means according to the cell type and placed in appropriate culture medium, again according to cell type. Viral particles are then contacted with the cells as described above, and the virus is allowed to transfect the cells. Cells can then be transplanted back into the subject's body, again by means standard for the cell type and tissue. If desired, prior to transplantation, the cells can be studied for degree of transfection by the virus, by known detection means and as described herein.

The present invention further provides a method of delivering a nucleic acid to a cell in a subject comprising administering to the subject an AAV2 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to a cell in the subject. Administration can be an ex vivo administration directly to a cell removed from a subject, such as any of the cells listed above, followed by replacement of the cell back into the subject, or administration can be in vivo administration to a cell in the subject. For ex vivo administration, cells are isolated from a subject by standard means according to the cell type and placed in appropriate culture medium, again according to cell type. Viral particles are then contacted with the cells as described above, and the virus is allowed to transfect the cells. Cells can then be transplanted back into the subject's body, again by means standard for the cell type and tissue. If desired, prior to transplantation, the cells can be studied for degree of transfection by the virus, by known detection means and as described herein.

Also provided is a method of delivering a nucleic acid to an epidermal cell in a subject comprising administering to the subject an AAV2 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to an epidermal cell in the subject.

In certain embodiments, the amino acid sequence that targets brain vascular endothelium targets brain vascular endothelium in a subject that has a disease, e.g., a lysosomal storage disease.

In certain embodiments, the amino acid sequence that targets brain vascular endothelium targets brain vascular endothelium in a subject that does not have a lysosomal storage disease.

In certain embodiments, the viral vector comprises a nucleic acid sequence encoding a therapeutic agent. In certain embodiments, the therapeutic agent is TPP1.

Certain embodiments of the present disclosure provide a cell comprising a viral vector as described herein.

In certain embodiments, the cell is a mammalian cell of a non-rodent mammal. In certain embodiments, the cell is a primate cell. In certain embodiments, the cell is a human cell. In certain embodiments, the cell is a non-human cell. In certain embodiments, the cell is in vitro. In certain embodiments, the cell is in vivo. In certain embodiments, the cell is an epidermal cell.

Certain embodiments of the present disclosure provide a method of treating a disease in a mammal comprising administering a viral vector or the cell as described herein to the mammal.

In certain embodiments, the mammal is human.

In certain embodiments, the disease is a lysosomal storage disease (LSD). In certain embodiments, the LSD is infantile or late infantile ceroid lipofuscinoses, Gaucher, Juvenile Batten, Fabry, MLD, Sanfilippo A, Late Infantile Batten, Hunter, Krabbe, Morquio, Pompe, Niemann-Pick C, Tay-Sachs, Hurler (MPS-I H), Sanfilippo B, Maroteaux-Lamy, Niemann-Pick A, Cystinosis, Hurler-Scheie (MPS-I H/S), Sly Syndrome (MPS VII), Scheie (MPS-I S), Infantile Batten, GM1 Gangliosidosis, Mucolipidosis type II/III, or Sandhoff disease.

In certain embodiments, the disease is a neurodegenerative disease. In certain embodiments, the neurodegenerative disease is Huntington's disease, ALS, hereditary spastic hemiplegia, primary lateral sclerosis, spinal muscular atrophy, Kennedy's disease, Alzheimer's disease, a polyglutamine repeat disease, or Parkinson's disease.

Certain embodiments of the present disclosure provide a method to deliver an agent to the central nervous system of a subject, comprising administering to the CSF with a viral vector described herein so that the transduced epidermal cells express the therapeutic agent and deliver the agent to the central nervous system of the subject. In certain embodiments, the viral vector transduces epidermal cells.
Certain embodiments of the present disclosure provide a viral vector or cell as described herein for use in medical treatments. Certain embodiments of the present disclosure provide a use of a viral vector or cell as described herein to prepare a medicament useful for treating a disease, e.g., a lysosomal storage disease, in a mammal. The vector may further comprise a lysosomal enzyme (e.g., a lysosomal hydrolase), a secreted protein, a nuclear protein, or a cytoplasmic protein. As used herein, the term “secreted protein” includes any secreted protein, whether naturally secreted or modified to contain a signal sequence so that it can be secreted.

Nucleic acid is “operably linked” when it is placed into a functional relationship with another nucleic acid sequence. Generally, “operably linked” means that the DNA sequences being linked are contiguous. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice. Additionally, multiple copies of the nucleic acid encoding enzymes may be linked together in the expression vector. Such multiple nucleic acids may be separated by linkers.

The present disclosure also provides a mammalian cell containing a vector described herein. The cell may be human, and may be from brain. The cell type may be a stem or progenitor cell population.

The present disclosure provides a method of treating a disease such as a genetic disease or cancer in a mammal by administered a polynucleotide, polypeptide, expression vector, or cell described herein. The genetic disease or cancer may be a lysosomal storage disease (LSD) such as infantile or late infantile ceroid lipofuscinosis, Gaucher, Juvenile Batten, Fabry, M.D., Sanfilippo A, Late Infantile Batten, Hunter, Krabbe, Morquio, Pompe, Niemann-Pick C, Tay-Sachs, Hunter (MPS-I H), Sanfilippo B, Maroteaux-Lamy, Niemann-Pick A, Cystinosis, Hunter-Scheie (MPS-I H/S), Sly Syndrome (MPS VII), Scheie (MPS-I S), Infantile Batten, GM1 Gangliosidosis, Mucolipidosis type II/III or Sandhoff disease.

The genetic disease may be a neurodegenerative disease, such as Huntington’s disease, ALS, hereditary spastic hemiplegia, primary lateral sclerosis, spinal muscular atrophy, Kennedy’s disease, Alzheimer’s disease, a polyglutamine repeat disease, or focal exposure such as Parkinson’s disease.

Certain aspects of the disclosure relate to polynucleotides, polypeptides, vectors, and genetically engineered cells (modified in vivo), and the use of them. In particular, the disclosure relates to a method for gene or protein therapy that is capable of both systemic delivery of a therapeutically effective dose of the therapeutic agent.

According to one aspect, a cell expression system for expressing a therapeutic agent in a mammalian recipient is provided. The expression system (also referred to herein as a “genetically modified cell”) comprises a cell and an expression vector for expressing the therapeutic agent. Expression vectors include, but are not limited to, viruses, plasmids, and other vehicles for delivering heterologous genetic material to cells. Accordingly, the term “expression vector” as used herein refers to a vehicle for delivering heterologous genetic material to a cell. In particular, the expression vector is a recombinant adenoviral, adeno-associated virus, or lentivirus or retrovirus vector.

The expression vector further includes a promoter for controlling transcription of the heterologous gene. The promoter may be an inducible promoter (described below). The expression system is suitable for administration to the mammalian recipient. The expression system may comprise a plurality of non-immortalized genetically modified cells, each cell containing at least one recombinant gene encoding at least one therapeutic agent.

The cell expression system can be formed in vivo. According to yet another aspect, a method for treating a mammalian recipient in vivo is provided. The method includes introducing an expression vector for expressing a heterologous gene product into a cell of the patient in situ, such as via intravenous administration. To form the expression system in vivo, an expression vector for expressing the therapeutic agent is introduced in vivo into the mammalian recipient i.e., where the vector migrates via the vasculature to the brain.

According to yet another aspect, a method for treating a mammalian recipient in vivo is provided. The method includes introducing the target protein into the patient in vivo.

The expression vector for expressing the heterologous gene may include an inducible promoter for controlling transcription of the heterologous gene product. Accordingly, delivery of the therapeutic agent in situ is controlled by exposing the cell in situ to conditions, which induce transcription of the heterologous gene.

The mammalian recipient may have a condition that is amenable to gene replacement therapy. As used herein, “gene replacement therapy” refers to administration to the recipient of exogenous genetic material encoding a therapeutic agent and subsequent expression of the administered genetic material in situ. Thus, the phrase “condition amenable to gene replacement therapy” embraces conditions such as genetic diseases (i.e., a disease condition that is attributable to one or more gene defects), acquired pathologies (i.e., a pathological condition which is not attributable to an inborn defect), cancers and prophylactic processes (i.e., prevention of a disease or of an undesired medical condition). Accordingly, as used herein, the term “therapeutic agent” refers to any agent or material, which has a beneficial effect on the mammalian recipient. Thus, “therapeutic agent” embraces both therapeutic and prophylactic molecules having nucleic acid or protein components.

According to one embodiment, the mammalian recipient has a genetic disease and the exogenous genetic material comprises a heterologous gene encoding a therapeutic agent for treating the disease. In yet another embodiment, the mammalian recipient has an acquired pathology and the exogenous genetic material comprises a heterologous gene encoding a therapeutic agent for treating the pathology. According to another embodiment, the patient has a cancer and the exogenous genetic material comprises a heterologous gene encoding an anti-neoplastic agent. In yet another embodiment, the patient has an undesired medical condition and the exogenous genetic material comprises a heterologous gene encoding a therapeutic agent for treating the condition.

As used herein, the term “lysosomal enzyme,” a “secreted protein,” a “nuclear protein,” or a “cytoplasmic protein” include variants or biologically active or inactive fragments of these polypeptides. A “variant” of one of the polypeptides is a polypeptide that is not completely identical to a native protein. Such variant protein can be obtained by altering the amino acid sequence by insertion, deletion or substitution of one or more amino acid. The amino acid sequence of the protein is modified, for example by substitution, to create a polypeptide having substantially the same or improved
qualities as compared to the native polypeptide. The substitu-
tion may be a conserved substitution. A "conserved sub-
stitution" is a substitution of an amino acid with another
amino acid having a similar side chain. A conserved sub-
stitution would be a substitution with an amino acid that makes
the smallest change possible in the charge of the amino acid
or size of the side chain of the amino acid (alternatively, in
the size, charge or kind of chemical group within the side
chain) such that the overall peptide retains its special con-
formation but has altered biological activity. For example,
commonly conserved changes might be Asp to Glu, Asn or
Gln; His to Lys, Arg or Phe; Asn to Gln, Asp or Glu and Ser
to Cys, Thr or Gly. Alanine is commonly used to substitute
for other amino acids. The 20 essential amino acids can be
grouped as follows: alanine, valine, leucine, isoleucine,
proline, phenylalanine, tryptophan and methionine having
nonpolar side chains; glycine, serine, threonine, cystine,
tyrosine, asparagine and glutamine having uncharged polar
side chains; aspartate and glutamate having acidic side
chains; and lysine, arginine, and histidine having basic side
chains.

The amino acid changes are achieved by changing the
codons of the corresponding nucleic acid sequence. It is
known that such polypeptides can be obtained based on
substituting certain amino acids for other amino acids in
the polypeptide structure in order to modify or improve biologi-
cal activity. For example, through substitution of alternative
amino acids, small conformational changes may be conferred
upon a polypeptide that results in increased activity.
Alternatively, amino acid substitutions in certain polypep-
tides may be used to provide residues, which may then be
linked to other molecules to provide peptide-molecule conjugates
which retain sufficient properties of the starting polypeptide
for other purposes.

One can use the hydrophobic index of amino acids in
confering interactive biological function on a polypeptide,
wherein it is found that certain amino acids may be substi-
tuted for other amino acids having similar hydrophobic
indices and still retain a similar biological activity. Alterna-
tively, substitution of like amino acids may be made on the
basis of hydrophilicity, particularly where the biological
function desired in the polypeptide to be generated in
intended for use in immunological embodiments. The greatest
local average hydrophilicity of a "protein", as governed
by the hydrophilicity of its adjacent amino acids, correlates
with its immunogenicity. Accordingly, it is noted that substi-
tutions can be made based on the hydrophilicity assigned
to each amino acid.

In using either the hydrophilicity index or hydrophobic
index, which assigns values to each amino acid, it is pref-
ered to conduct substitutions of amino acids where these
values are ±2, with ±1 being particularly preferred, and those
with in ±0.5 being the most preferred substitutions.

The variant protein has at least 50%, at least about 80%,
or even at least about 90% but less than 100%, contiguous
amino acid sequence homology or identity to the amino acid
sequence of a corresponding native protein.

The amino acid sequence of the variant polypeptide
Corresponds essentially to the native polypeptide's amino
acid sequence. As used herein "correspond essentially to"
refers to a polypeptide sequence that will elicit a biological
response substantially the same as the response generated by
the native protein. Such a response may be at least 60% of
the level generated by the native protein, and may even be
at least 80% of the level generated by native protein.

A variant may include amino acid residues not present in
the corresponding native protein or deletions relative to the

A variant may also be a truncated "fragment" as compared to the corresponding
native protein, i.e., only a portion of a full-length protein. Protein variants also include peptides having at least one
D-amino acid.

The variant protein may be expressed from an isolated
dNA sequence encoding the variant protein. "Recombinant"
is defined as a peptide or nucleic acid produced by the
processes of genetic engineering. It should be noted that it
is well-known in the art that, due to the redundancy in the
gene code, individual nucleotides can be readily
exchanged in a codon, and still result in an identical amino
acid sequence. The terms "protein," "peptide" and "polypep-
tide" are used interchangeably herein.

The present disclosure provides methods of treating a
disease in a mammal by administering an expression vector
to a cell or patient. For the gene therapy methods, a person
having ordinary skill in the art of molecular biology and
gene therapy would be able to determine, without undue
experimentation, the appropriate dosages and routes of
administration of the expression vector used in the novel
methods of the present disclosure.

According to one embodiment, the cells are transformed
or otherwise genetically modified in vivo. The cells from the
mammalian recipient are transformed (i.e., transduced or
transfected) in vivo with a vector containing exogenous
material for expressing a heterologous (e.g., recombi-

nant) gene encoding a therapeutic agent and the therapeu-
tic agent is delivered in situ.

As used herein, "exogenous genetic material" refers to a
nucleic acid or an oligonucleotide, either natural or syn-
thetic, that is not naturally found in the cells; or if it is
naturally found in the cells, it is not transcribed or expressed
at biologically significant levels by the cells. Thus, "exog-
eneous genetic material" includes, for example, a non-natu-
rlly occurring nucleic acid that can be transcribed into
anti-sense RNA, as well as a "heterologous gene" (i.e., a
gene encoding a protein which is not expressed or is
expressed at biologically insignificant levels in a naturally-
occurring cell of the same type).

In the certain embodiments, the mammalian recipient has
a condition that is amenable to gene replacement therapy.
As used herein, "gene replacement therapy" refers to admin-
istration to the recipient of exogenous genetic material encod-
ing a therapeutic agent and subsequent expression of the
administered genetic material in situ. Thus, the phrase
"condition amenable to gene replacement therapy"
embraces conditions such as genetic diseases (i.e., a disease
condition that is attributable to one or more gene defects),
aquired pathologies (i.e., a pathological condition which is
not attributable to an inborn defect), cancers and prophy-
lactic processes (i.e., prevention of a disease or of an
undesired medical condition). Accordingly, as used herein,
the term "therapeutic agent" refers to any agent or material,
which has a beneficial effect on the mammalian recipient.
Thus, "therapeutic agent" embraces both therapeutic and
prophylactic molecules having nucleic acid (e.g., antisense
RNA) and/or protein components.

Alternatively, the condition amenable to gene replace-
ment therapy is a prophylactic process, i.e., a process for
preventing disease or an undesired medical condition. Thus,
the instant disclosure embraces a cell expression system for
delivering a therapeutic agent that has a prophylactic func-
tion (i.e., a prophylactic agent) to the mammalian recipient.

In summary, the term "therapeutic agent" includes, but is
not limited to, agents associated with the conditions listed
above, as well as their functional equivalents. As used
herein, the term "functional equivalent" refers to a molecule (e.g., a peptide or protein) that has the same or an improved beneficial effect on the mammalian recipient as the therapeutic agent of which it is deemed a functional equivalent. The above-disclosed therapeutic agents and conditions amenable to gene replacement therapy are merely illustrative and are not intended to limit the scope of the instant disclosure. The selection of a suitable therapeutic agent for treating a known condition is deemed to be within the scope of one of ordinary skill in the art without undue experimentation.

AAV2 Vectors

In one embodiment, a viral vector of the disclosure is an AAV2 vector. An "AAV2" vector refers to an adenov-associated virus, and may be used to refer to the naturally occurring wild-type virus itself or derivatives thereof. The term covers all subtypes, serotypes and pseudotypes, and both naturally occurring and recombinant forms, except where required otherwise. As used herein, the term "serotype" refers to an AAV which is identified by and distinguished from other AAV's based on capsid protein reactivity with defined antisera, e.g., there are eight known serotypes of primate AAVs, AAV-1 to AAV-8. For example, serotype AAV2 is used to refer to an AAV which contains capsid proteins encoded from the cap gene of AAV2 and a genome containing 5' and 3' ITR sequences from the same AAV2 serotype. As used herein, for example, rAAV1 may be used to refer to an AAV having both capsid proteins and 5'-3' ITRs from the same serotype or it may refer to an AAV having capsid proteins from one serotype and 5'-3' ITRs from a different AAV serotype, e.g., capsid from AAV serotype 2 and ITRs from AAV serotype 5. For each example illustrated herein the description of the vector design and production describes the serotype of the capsid and 5'-3' ITR sequences. The abbreviation "rAAV" refers to recombinant adenoviral vector, also referred to as a recombinant AAV vector (or "rAAV vector").

An "AAV virus" or "AAV viral particle" refers to a viral particle composed of at least one AAV capsid protein (preferably all of the capsid proteins of a wild-type AAV) and an encapsidated polynucleotide. If the particle comprises heterologous polynucleotide (i.e., a polynucleotide other than a wild-type AAV genome such as a transferrase to be delivered to a mammalian cell), it is typically referred to as "rAAV".

In one embodiment, the AAV expression vectors are constructed using known techniques to at least provide as operatively linked components in the direction of transcription, control elements including a transcriptional initiation region, the DNA of interest and a transcriptional termination region. The control elements are selected to be functional in a mammalian cell. The resulting construct which contains the operatively linked components is flanked (5' and 3') with functional AAV ITR sequences.

By "adenov-associated virus inverted terminal repeats" or "AAV ITRs" is meant the art-recognized regions found at each end of the AAV genome which function together in cis as origins of DNA replication and as packaging signals for the virus. AAV ITRs, together with the AAV rep coding region, provide for the efficient excision and rescue from, and integration of a nucleotide sequence interposed between two flanking ITRs into a mammalian cell genome.

The nucleotide sequences of AAV ITR regions are known. As used herein, an "AAV ITR" need not have the wild-type nucleotide sequence depicted, but may be altered, e.g., by the insertion, deletion or substitution of nucleotides. Additionally, the AAV ITR may be derived from any of several AAV serotypes, including without limitation, AAV1, AAV2, AAV3, AAV4, AAV5, AAV7, etc. Furthermore, 5' and 3' ITRs which flank a selected nucleotide sequence in an AAV vector need not necessarily be identical or derived from the same AAV serotype or isolate, so long as they function as intended, i.e., to allow for excision and rescue of the sequence of interest from a host cell genome or vector, and to allow integration of the heterologous sequence into the recipient cell genome when AAV Rep gene products are present in the cell.

In one embodiment, AAV ITRs can be derived from any of several AAV serotypes, including without limitation, AAV1, AAV2, AAV3, AAV4, AAV5, AAV7, etc. Furthermore, 5' and 3' ITRs which flank a selected nucleotide sequence in an AAV expression vector need not necessarily be identical or derived from the same AAV serotype or isolate, so long as they function as intended, i.e., to allow for excision and rescue of the sequence of interest from a host cell genome or vector, and to allow integration of the DNA molecule into the recipient cell genome when AAV Rep gene products are present in the cell.

In one embodiment, AAV capsids can be derived from AAV2. Suitable DNA molecules for use in AAV vectors will be less than about 5 kilobases (kb), less than about 4.5 kb, less than about 4 kb, less than about 3.5 kb, less than about 3 kb, less than about 2.5 kb in size and are known in the art.

In one embodiment, theselected nucleotide sequence is operably linked to control elements that direct the transcription or expression thereof in the subject in vivo. Such control elements can comprise control sequences normally associated with the selected gene. Alternatively, heterologous control sequences can be employed. Useful heterologous control sequences generally include those derived from sequences encoding mammalian or viral genes. Examples include, but are not limited to, the SV40 early promoter, mouse mammary tumor virus (LTR) promoter, adeovirus major late promoter (Ad MLP); a herpes simplex virus (HSV) promoter, a cytomegalovirus (CMV) promoter such as the CMV immediate early promoter region (CMVIE), a rous sarcoma virus (RSV) promoter, pol II promoters, pol III promoters, synthetic promoters, hybrid promoters, and the like. In addition, sequences derived from nonviral genes, such as the murine metallothionein gene, will also find use herein. Such promoter sequences are commercially available from, e.g., Stratagene (San Diego, Calif.).

In one embodiment, both heterologous promoters and other control elements, such as CNS-specific and inducible promoters, enhancers and the like, will be of particular use. Examples of heterologous promoters include the CMV promoter. Examples of CNS-specific promoters include those isolated from the genes from myelin basic protein (MBP), glial fibrillary acid protein (GFAP), and neuron specific enolase (NSE). Examples of inducible promoters include DNA responsive elements for ceddyson, tetracycline, hypoxia and auxin.

In one embodiment, the AAV expression vector which harbors the DNA molecule of interest bounded by AAV ITRs, can be constructed by directly inserting the selected sequence(s) into an AAV genome which has had the major AAV open reading frames ("ORFs") excised therefrom. Other portions of the AAV genome can also be deleted, so long as a sufficient portion of the ITRs remain to allow for replication and packaging functions. Such constructs can be designed using techniques well known in the art.

Alternatively, AAV ITRs can be excised from the viral genome or from an AAV vector containing the same and fused 5' and 3' of a selected nucleic acid construct that is
present in another vector using standard ligation techniques. For example, ligation can be accomplished in 20 mM Tris-Cl pH 7.5, 10 mM MgCl₂, 10 mM DTT, 33 μg/ml BSA, 10 mM-50 mM NaCl, and either 40 nM ATP, 0.01-0.02 (Weiss) units T4 DNA ligase at 0° C. (for “sticky end” ligation) or 1 nM ATP, 0.5-0.6 (Weiss) units T4 DNA ligase at 14° C. (for “blunt end” ligation). Intermolecular “sticky end” ligations are usually performed at 30-100 μg/ml total DNA concentrations (5-100 nM total end concentration). AAV vectors which contain ITRs.

Additionally, chimeric genes can be produced synthetically to include AAV ITR sequences arranged 5' and 3' of one or more selected nucleic acid sequences. Preferred codons for expression of the chimeric gene sequence in mammalian CNS cells can be used. The complete chimeric sequence is assembled from overlapping oligonucleotides prepared by standard methods.

In order to produce rAAV virions, an AAV expression vector is introduced into a suitable host cell using known techniques, such as transfection. A number of transfection techniques are generally known in the art. See, e.g., Sambrook et al. (1989) Molecular Cloning, a laboratory manual, Cold Spring Harbor Laboratories, New York. Particularly suitable transfection methods include calcium phosphate co-precipitation, direct micro-injection into cultured cells, electroporation, liposome mediated gene transfer, lipid-mediated transduction, and nucleic acid delivery using high velocity microprojectiles.

In one embodiment, suitable host cells for producing rAAV virions include microorganisms, yeast cells, insect cells, and mammalian cells, that can be, or have been, used as recipients of a heterologous DNA molecule. The term includes the progeny of the original cell which has been transfected. Thus, a “host cell” as used herein generally refers to a cell which has been transfected with an exogenous DNA sequence. Cells from the stable human cell line, 293 (readily available through, e.g., the American Type Culture Collection under Accession Number ATCC CRL1573) can be used in the practice of the present disclosure. Particularly, the human cell line 293 is a human embryonic kidney cell line that has been transfected with adenovirus type-5 DNA fragments, and expresses the adenoviral E1a and E1b genes. The 293 cell line is readily transfected, and provides a particularly convenient platform in which to produce rAAV virions.

By “AAV rep coding region” is meant the art-recognized region of the AAV genome which encodes the replication proteins Rep 78, Rep 68, Rep 52 and Rep 40. These Rep expression products have been shown to possess many functions, including recognition, binding and nicking of the AAV origin of DNA replication, DNA helicase activity and modulation of transcription from AAV (or other heterologous) promoters. The Rep expression products are collectively required for replicating the AAV genome. Suitable homologues of the AAV rep coding region include the human herpesvirus 6 (HHV-6) rep gene which is also known to mediate AAV-2 DNA replication.

By “AAV cap coding region” is meant the art-recognized region of the AAV genome which encodes the capsid proteins VP1, VP2, and VP3, or functional homologues thereof. These Cap expression products supply the packaging functions which are collectively required for packaging the viral genome.

In one embodiment, AAV helper functions are introduced into the host cell by transfecting the host cell with an AAV helper construct either prior to, or concurrently with, the transfection of the AAV expression vector. AAV helper constructs are thus used to provide at least transient expression of AAV rep and/or cap genes to complement missing AAV functions that are necessary for productive AAV infection. AAV helper constructs lack AAV ITRs and can neither replicate nor package themselves. These constructs can be in the form of a plasmid, phage, transposon, cosmid, virus, or virion. A number of AAV helper constructs have been described, such as the commonly used plasmids pAAV/Ad and pM29+45 which encode both Rep and Cap expression products. A number of other vectors have been described which encode Rep and/or Cap expression products.

Methods of delivery of viral vectors include injecting the AAV2 into the CNS. Generally, rAAV virions may be introduced into cells of the CNS using either in vivo or in vitro transduction techniques. If transduced in vitro, the desired recipient cell will be removed from the subject, transduced with rAAV virions and reintroduced into the subject. Alternatively, syngeneic or xenogeneic cells can be used where those cells will not generate an inappropriate immune response in the subject.

Suitable methods for the delivery and introduction of transduced cells into a subject have been described. For example, cells can be transduced in vitro by combining recombinant AAV virions with CNS cells e.g., in appropriate media, and screening for those cells harboring the DNA of interest can be screened using conventional techniques such as Southern blots and/or PCR, or by using selectable markers. Transduced cells can then be formulated into pharmaceutical compositions, described more fully below, and the composition introduced into the subject by various techniques, such as by grafting, intramuscular, intravenous, subcutaneous and intraperitoneal injection.

In one embodiment, pharmaceutical compositions will comprise sufficient genetic material to produce a therapeutically effective amount of the nucleic acid of interest, i.e., an amount sufficient to reduce or ameliorate symptoms of the disease state in question or an amount sufficient to confer the desired benefit. The pharmaceutical compositions will also contain a pharmaceutically acceptable excipient. Such excipients include any pharmaceutical agent that does not itself induce the production of antibodies harmful to the individual receiving the composition, and which may be administered without undue toxicity. Pharmaceutically acceptable excipients include, but are not limited to, sorbitol, Tween80, and liquids such as water, saline, glycerol and ethanol. Pharmaceutically acceptable salts can be included therein, for example, mineral acid salts such as hydrochlorides, hydrobromides, phosphates, sulfates, and the like; and the salts of organic acids such as acetates, propionates, malonates, benzoates, and the like. Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present in such vehicles. A thorough discussion of pharmaceutically acceptable excipients is available in Remington’s Pharmaceutical Sciences (Mack Pub. Co., N.J. 1991).

As is apparent to those skilled in the art in view of the teachings of this specification, an effective amount of viral vector which must be added can be empirically determined. Administration can be effected in one dose, continuously or intermittently throughout the course of treatment. Methods of determining the most effective means and dosages of administration are well known to those of skill in the art and will vary with the viral vector, the composition of the therapy, the target cells, and the subject being treated. Single and multiple administrations can be carried out with the dose level and pattern being selected by the treating physician.
It should be understood that more than one transgene could be expressed by the delivered viral vector. Alternatively, separate vectors, each expressing one or more different transgenes, can also be delivered to the CNS as described herein. Furthermore, it is also intended that the viral vectors delivered by the methods of the present disclosure be combined with other suitable compositions and therapies.

Methods for Introducing Genetic Material into Cells

The exogenous genetic material (e.g., a DNA encoding one or more therapeutic proteins) is introduced into the cell ex vivo or in vivo by genetic transfer methods, such as transfection or transduction, to provide a genetically modified cell. Various expression vectors (i.e., vehicles for facilitating delivery of exogenous genetic material into a target cell) are known to one of ordinary skill in the art.

As used herein, “transfection of cells” refers to the acquisition by a cell of new genetic material by incorporation of added DNA. Thus, transfection refers to the insertion of nucleic acid into a cell using physical or chemical methods. Several transfection techniques are known to those of ordinary skill in the art including: calcium phosphate DNA co-precipitation; DEAE-dextran; electroporation; cationic liposome-mediated transfection; and tungsten particle-facilitated microparticle bombardment. Strontium phosphate DNA co-precipitation is another possible transfection method.

In contrast, “transduction of cells” refers to the process of transferring nucleic acid into a cell using a DNA or RNA virus. A RNA virus (i.e., a retrovirus) for transferring a nucleic acid into a cell is referred to herein as a transducing chimeric retrovirus. Exogenous genetic material contained within the retrovirus is incorporated into the genome of the transduced cell. A cell that has been transduced with a chimeric DNA virus (e.g., an adenovirus carrying a DNA encoding a therapeutic agent), will not have the exogenous genetic material incorporated into its genome but will be capable of expressing the exogenous genetic material that is maintained extrachromosomally within the cell.

Typically, the exogenous genetic material includes the heterologous gene (usually in the form of a CDNA comprising the exons encoding for the therapeutic protein) together with a promoter to control transcription of the new gene. The promoter characteristically has a specific nucleotide sequence necessary to initiate transcription. Optionally, the exogenous genetic material further includes additional sequences (i.e., enhancers) required to obtain the desired gene transcription activity. For the purpose of this discussion an “enhancer” is simply any non-translated DNA sequence which works contiguous with the coding sequence (in cis) to change the basal transcription level dictated by the promoter. The exogenous genetic material may be introduced into the cell genome immediately downstream from the promoter so that the promoter and coding sequence are operatively linked so as to permit transcription of the coding sequence. A retroviral expression vector may include an exogenous promoter element to control transcription of the inserted exogenous gene. Such exogenous promoters include both constitutive and inducible promoters.

Naturally occurring constitutive promoters control the expression of essential cell functions. As a result, a gene under the control of a constitutive promoter is expressed under all conditions of cell growth. Exemplary constitutive promoters include the promoters for the following genes which encode certain constitutive or “housekeeping” functions: hypoxanthine phosphoribosyl transferase (HPRT), dihydrofolate reductase (DHFR), adenosine deaminase, phosphoglycerol kinase (PGK), pyruvate kinase, phosphoglycerol mutase, the actin promoter, and other constitutive promoters known to those of skill in the art. In addition, many viral promoters function constitutively in eukaryotic cells. These include: the early and late promoters of SV40; the long terminal repeats (LTRs) of Moloney Leukemia Virus and other retroviruses; and the thymidine kinase promoter of Herpes Simplex Virus, among many others. Accordingly, any of the above-referenced constitutive promoters can be used to control transcription of a heterologous gene insert.

Genes that are under the control of inducible promoters are expressed only or to a greater degree, in the presence of an inducing agent, (e.g., transcription under control of the metallothionein promoter is greatly increased in presence of certain metal ions). Inducible promoters include responsive elements (REs) which stimulate transcription when their inducing factors are bound. For example, these REs for serum factors, steroid hormones, retinoic acid and cyclic AMP. Promoters containing a particular RE can be chosen in order to obtain an inducible response and in some cases, the RE itself may be attached to a different promoter, thereby conferring inducibility to the recombinant gene. Thus, by selecting the appropriate promoter (constitutive versus inducible; strong versus weak), it is possible to control both the existence and level of expression of a therapeutic agent in the genetically modified cell. If the gene encoding the therapeutic agent is under the control of an inducible promoter, delivery of the therapeutic agent in situ is triggered by exposing the genetically modified cell in situ to conditions for permitting transcription of the therapeutic agent, e.g., by intraperitoneal injection of specific inducers of the inducible promoters which control transcription of the agent.

For example, in situ expression by genetically modified cells of a therapeutic agent encoded by a gene under the control of the metallothionein promoter, is enhanced by contacting the genetically modified cells with a solution containing the appropriate (i.e., inducing) metal ions in situ.

Accordingly, the amount of therapeutic agent that is delivered in situ is regulated by controlling such factors as: (1) the nature of the promoter used to direct transcription of the inserted gene, (i.e., whether the promoter is constitutive or inducible, strong or weak); (2) the number of copies of the exogenous gene that are inserted into the cell; (3) the number of transduced/transfected cells that are administered (e.g., implanted) to the patient; (4) the size of the implant (e.g., graft or encapsulated expression system); (5) the number of implants; (6) the length of time the transduced/transfected cells or implants are left in place; and (7) the production rate of the therapeutic agent by the genetically modified cell. Selection and optimization of these factors for delivery of a therapeutically effective dose of a particular therapeutic agent is deemed to be within the scope of one of ordinary skill in the art without undue experimentation, taking into account the above-disclosed factors and the clinical profile of the patient.

In addition to at least one promoter and at least one heterologous nucleic acid encoding the therapeutic agent, the expression vector may include a selection gene, for example, a neomycin resistance gene, for facilitating selection of cells that have been transfected or transduced with the expression vector. Alternatively, the cells are transfected with two or more expression vectors, at least one vector containing the gene(s) encoding the therapeutic agent(s), the other vector containing a selection gene. The selection of a suitable promoter, enhancer, selection gene and/or signal
sequence (described below) is deemed to be within the scope of one of ordinary skill in the art without undue experimentation.

The therapeutic agent can be targeted for delivery to an extracellular, intracellular, or membrane location. If it is desirable for the gene product to be secreted from the cells, the expression vector is designed to include an appropriate secretion “signal” sequence for secreting the therapeutic gene product from the cell to the extracellular milieu. If it is desirable for the gene product to be retained within the cell, this secretion signal sequence is omitted. In a similar manner, the expression vector can be constructed to include “retention” signal sequences for anchoring the therapeutic agent within the cell plasma membrane. For example, all membrane proteins have hydrophobic transmembrane regions, which stop translocation of the protein in the membrane and do not allow the protein to be secreted. The construction of an expression vector including signal sequences for targeting a gene product to a particular location is deemed to be within the scope of one of ordinary skill in the art without the need for undue experimentation.

EXAMPLE 1

Treating Central Nervous System Disorders Via Cerebral Spinal Fluid (CSF) in Large Mammals

Lysosomal storage disorders (LSDs) constitute a large class of inherited metabolic disorders. Most LSDs are caused by lysosomal enzyme deficiencies which lead to organ damage and often central nervous system (CNS) degeneration. Late infantile neuronal ceroid lipofuscinosis (LINCL) is an autosomal recessive neurodegenerative disease caused by mutations in CLN2, which encodes the lysosomal protease tripeptidyl peptidase 1 (TPP1). LINCL is characterized clinically by progressive motor and cognitive decline, and premature death. Enzyme-replacement therapy (ERT) is currently available for lysosomal storage diseases affecting peripheral tissues, but has not been used in patients with central nervous system (CNS) involvement. A recent study investigated whether enzyme delivery through the cerebrospinal fluid was a potential alternative route to the CNS for LINCL (Chang et al., Molecular Therapy 16:649-656, 2008). In this study, the investigators tested if intraventricular delivery of TPP1 to the LINCL mouse was efficacious. They found that infusion of recombinant human TPP1 through an intraventricular cannula led to enzyme distribution in several regions of the brain of treated mice. In vitro activity assays confirmed increased TPP1 activity throughout the rostral-caudal extent of the brain. Treated mice showed attenuated neuropathology, and decreased resting tremor relative to vehicle-treated mice.

The next step was to determine whether long-term, steady-state levels of therapeutic enzymes could be achieved in a mammal. It was discovered that ependymal cells (cells that line the ventricles in the brain) can be transduced and secrete a targeted enzyme into the cerebral spinal fluid (CSF). It was determined that adeno-associated virus (AAV4) can transduce the ependyma in a mouse model with high efficiency. Davidson et al., PNAS. 28:3428-3432, 2000. It was found that in mice there was a normalization of stored substrate levels in disease brain after AAV4 treatment.

In the present work, it was investigated whether global delivery of a vector could be effectively performed in order to achieve steady-state levels of enzyme in the CSF. First, a vector needed to be found that could transduce ependymal cells (cells that line the ventricles) in the brain of larger mammals. Studies were performed in a dog model of LINCL and a non-human primate model of LINCL. The LINCL dogs are normal at birth, but develop neurological signs around 7 months, testable cognitive deficits at ~5-6 months, seizures at 10-11 months, and progressive visual loss.

An adeno-associated virus (AAV) was selected as the vector because of its small size (20 nm), most of its genetic material can be removed ("gutted") so that no viral genes are present, and so that it is replication incompetent. It was previously tested whether adeno-associated virus type 4 (AAV4) vectors could mediate global functional and pathological improvements in a murine model of mucopolysaccharidosis type VII (MPS VII) caused by beta-glucuronidase deficiency (Liu et al., J. Neuroscience, 25(41):9321-9327, 2005). Recombinant AAV vectors encoding beta-glucuronidase were injected unilaterally into the lateral ventricle of MPS VII mice with established disease. Transduced ependyma expressed high levels of recombinant enzyme, with secreted enzyme penetrating cerebral and cerebellar structures, as well as the brainstem. Immunohistochemical studies revealed close association of recombinant enzyme and brain microvasculature, indicating that beta-glucuronidase reached brain parenchyma via the perivascular spaces lining blood vessels. Aversive associative learning was tested by context fear conditioning. Compared with age-matched heterozygous controls, affected mice showed impaired conditioned fear response and context discrimination. This behavioral deficit was reversed 6 weeks after gene transfer in AAV4 beta-glucuronidase-treated MPS VII mice. The data show that ependymal cells can serve as a source of enzyme secretion into the surrounding brain parenchyma and CSF.

Surprisingly, however, when these studies were extended to large mammals (i.e., dogs and non-human primates), the AAV4 vectors were not effective in targeting the ependyma in these animals. Instead, an AAV2 vector needed to be used. Results of these experiments are shown for dogs (FIG. 1) and nonhuman primates (NHP, FIG. 2). Briefly, rAAV2 was generated encoding TPP1 (AAV2-CLN2), and injected intraventricularly to transduce ependyma (Liu et al., J. Neuroscience, 25(41):9321-9327, 2005). TPP1 is the enzyme deficient in LINCL. The data indicated that ependymal transduction in NHP brain resulted in a significant increase of enzyme in CSF (FIG. 3). The results indicated elevated levels of TPP1 activity in various brain regions, where the vertical axis show % control of activity (FIG. 4).

In the first dog that was treated, the delivery of vector was suboptimal, but still exhibited CLN2 activity in the brain. Subsequent dogs underwent IVC delivery with stereotaxy. It was found that the cognitive abilities of the treated dogs were significantly improved over a non-treated dog, as measured by T-maze performance (FIG. 5). Further, the effects of IVC delivery of AAV2-CLN2 in the dog model of LINCL were very pronounced. In the untreated (−/−) animal, large ventricles are present, whereas the brains of the untreated control and the treated animals did not exhibit ventricles. Following delivery of AAV-TPP1 to ventricles of LINCL dogs, detectable enzyme activity was noted in various brain regions, including the cerebellum and upper spinal cord. In two living additional affected dogs, brain atrophy was significantly attenuated, longevity was increased and cognitive function was improved. Finally, in NHP, we show that this method can achieve TPP1 activity levels 2-5 fold above wildtype.

Several AAV vectors were generated and tested to determine the optimal combination of ITR and capsid. Five different combinations were produced, once it was deter-
minded that the AAV2 ITR was most effective: AAV2/1 (i.e., AAV2 ITR and AAV1 capsid), AAV2/2, AAV2/4, AAV2/5, and AAV2/8. It was discovered that AAV2/2 worked much better in the large mammals (dogs and NHPI), followed by AAV2/8, AAV2/5, AAV2/1 and AAV2/4. This was quite surprising because the order of effectiveness of the viral vectors is the opposite of what was observed in mice.

Thus, the present work has shown that ventricular lining cells can be a source of recombinant enzyme in CSF for distribution throughout the brain, and that AAV2/2 is an effective vehicle for administering therapeutic agents, such as the gene encoding C1N2 (TPP1) in dogs and nonhuman primates.

All publications, patents and patent applications are incorporated herein by reference. While in the foregoing specification this invention has been described in relation to certain preferred embodiments thereof, and many details have been set forth for purposes of illustration, it will be apparent to those skilled in the art that the invention is susceptible to additional embodiments and that certain of the details described herein may be varied considerably without departing from the basic principles of the invention.

The use of the terms "a" and "an" and "the" and similar referents in the context of describing the invention are to be construed to cover both the singular and the plural, unless otherwise indicated herein or clearly contradicted by context. The terms "comprising," "having," "including," and "containing" are to be construed as open-ended terms (i.e., meaning "including, but not limited to") unless otherwise noted. Recitation of ranges of values herein are merely intended to serve as a shorthand method of referring individually to each separate value falling within the range, unless otherwise indicated herein, and each separate value is incorporated into the specification as if it were individually recited herein. All methods described herein can be performed in any suitable order unless otherwise indicated herein or otherwise clearly contradicted by context. The use of any and all examples, or exemplary language (e.g., "such as") provided herein, is intended merely to better illuminate the invention and does not pose a limitation on the scope of the invention unless otherwise claimed. No language in the specification should be construed as indicating any non-claimed element as essential to the practice of the invention.

Embodiments of this invention are described herein, including the best mode known to the inventors for carrying out the invention. Variations of those embodiments may become apparent to those of ordinary skill in the art upon reading the foregoing description. The inventors expect skilled artisans to employ such variations as appropriate, and the inventors intend for the invention to be practiced otherwise than as specifically described herein. Accordingly, this invention includes all modifications and equivalents of the subject matter recited in the claims appended hereto as permitted by applicable law. Moreover, any combination of the above-described elements in all possible variations thereof is encompassed by the invention unless otherwise indicated herein or otherwise clearly contradicted by context.

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What is claimed is:

1. A method of treating a central nervous system disease in a non-rodent mammal comprising:
   delivering a therapeutic protein to cerebrospinal fluid (CSF) of the non-rodent mammal by a method
   comprising,
   (a) intravenicularly administering to the cerebrospinal fluid (CSF) of the non-rodent mammal an AAV2
   particle comprising an AAV2 capsid protein and a vector comprising a nucleic acid encoding the
   therapeutic protein inserted between a pair of AAV inverted terminal repeats in a manner effective to
   infect an ependymal cell in the non-rodent mammal, wherein the ependymal cell secretes the therapeutic protein so as to
   treat the central nervous system disease; or
   (b) intravenicularly administering to ependymal cells of the mammal an AAV2 particle comprising an AAV2
   capsid protein and a vector comprising a nucleic acid encoding the therapeutic protein inserted between a
   pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to the ependymal cell, wherein
   the ependymal cell secretes the therapeutic protein so as to treat the central nervous system disease;
   wherein the central nervous system disease is a lysosomal storage disease (LSD) and the therapeutic protein is a
   lysosomal hydrolase.

2. A method of delivering a lysosomal hydrolase protein to the central nervous system of a non-rodent mammal having a lysosomal storage disease (LSD), comprising intravenicularly administering to the cerebrospinal fluid (CSF) of the non-rodent mammal an AAV2 particle comprising an AAV2 capsid protein and a vector comprising a nucleic acid encoding the protein lysosomal hydrolase inserted between a pair of AAV inverted terminal repeats in a manner effective to infect ependymal cells in the non-rodent mammal, wherein such that the ependymal cells secrete the protein lysosomal hydrolase into the CSF of the non-rodent mammal and deliver the lysosomal hydrolase to the central nervous system of the mammal.

3. The method of claim 1, wherein the non-rodent mammal is a primate or dog.

4. The method of claim 3, wherein the non-rodent mammal is a primate.

5. The method of claim 3, wherein the non-rodent mammal is a dog.

6. The method of claim 1 or 2, wherein the protein LSD is late infantile ceroid lipofuscinosis (LINCL) and the lysosomal hydrolase is tripeptidyl peptidase 1 (TPP1).

7. The method of claim 1, wherein the non-rodent mammal is human.

8. The method of claim 1 or 2, wherein the LSD is infantile or late infantile ceroid lipofuscinosis (LINCL), neuroopathic Gaucher, Juvenile Batten, Fabry, MLND, Sandhoff, Hunter, Krabbe, Morquio, Pompe, Niemann-Pick C, Tay-Sachs, Hurler (MPS-I H), Sanfilippo B, Maroteaux-Lamy, Niemann-Pick A, Cystinosis, Hurler-Scheie (MPS-I H/S), Sly Syndrome (MPS VII), Scheie (MPS-I S), Infantile Batten, GM1 Gangliosidosis, Mucolipidosis type II/III, or Sandhoff disease.

9. The method of claim 8, wherein the LSD is late infantile ceroid lipofuscinoses (LINCL).

10. The method of claim 8, wherein the LSD is Juvenile Batten or Infantile Batten disease.

11. The method of claim 1, wherein at least one of the pair of AAV inverted terminal repeats is an AAV2 ITR.

12. The method of claim 2, wherein at least one of the pair of AAV inverted terminal repeats is an AAV2 ITR.

13. The method of claim 1, wherein the ependymal cells provide a source of secreted protein that is distributed to or penetrates the cerebral or cerebellar structures of the non-rodent mammal.

14. The method of claim 1, wherein the ependymal cells provide a source of secreted protein that is distributed to or penetrates the upper spinal cord or brainstem of the non-rodent mammal.

15. The method of claim 1, wherein the ependymal cells provide a source of secreted protein that is distributed throughout the brain of the non-rodent mammal.

16. The method of claim 10, wherein the lysosomal hydrolase is TPP1 and the non-rodent mammal is a human.

17. The method of claim 2, wherein the non-rodent mammal is a primate or dog.

18. The method of claim 17, wherein the non-rodent mammal is a primate.

19. The method of claim 18, wherein the primate is human.

20. The method of claim 2, wherein the non-rodent mammal is a dog.

21. The method of claim 2, wherein the lysosomal hydrolase is TPP1.

22. The method of claim 2, wherein the ependymal cells provide a source of secreted protein that is distributed to or penetrates the cerebral or cerebellar structures of the non-rodent mammal.

23. A method of treating a central nervous system disease in a non-rodent mammal comprising:
   delivering a therapeutic protein to cerebrospinal fluid (CSF) of the non-rodent mammal by a method 
   comprising,
   (a) intravenicularly administering to the cerebrospinal fluid (CSF) of the non-rodent mammal an AAV2 
   particle comprising an AAV2 capsid protein and a vector comprising a nucleic acid encoding the 
   therapeutic protein inserted between a pair of AAV inverted terminal repeats in a manner effective to 
   infect an ependymal cell in the non-rodent mammal, wherein the ependymal cell secretes the therapeutic protein so as to 
   treat the central nervous system disease; or
   (b) intravenicularly administering to ependymal cells of the mammal an AAV2 particle comprising an AAV2 
   capsid protein and a vector comprising a nucleic acid encoding the therapeutic protein inserted between a 
   pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to the ependymal cell, wherein
   the ependymal cell secretes the therapeutic protein so as to treat the central nervous system disease;
   wherein the central nervous system disease is a lysosomal storage disease (LSD) and the therapeutic protein is a 
   lysosomal hydrolase.
23. The method of claim 2, wherein the ependymal cells provide a source of secreted protein that is distributed to or penetrates the upper spinal cord or brainstem of the non-rodent mammal.

24. The method of claim 2, wherein the ependymal cells provide a source of secreted protein that is distributed throughout the brain of the non-rodent mammal.