Gene Therapy Corrects Brain and Behavioral Pathologies in CLN6-Batten Disease

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INTRODUCTION

Batten disease (also called neuronal ceroid lipofuscinosis [NCLs]) comprises a family of rare, genetic, lysosomal storage disorders that are universally fatal. This family of diseases is caused by mutations in one of approximately 13 known ceroid-lipofuscinosis neuronal (CLN)-related genes.1 Collectively, Batten disease is the most prevalent cause of neurodegenerative disease in children, with an incidence of 2–4/100,000 live births.2,3 Common disease symptomology and pathology in this family of diseases involves progressive visual, motor, and mental deterioration, with shortened lifespan.

Gene Therapy Corrects Brain and Behavioral Pathologies in CLN6-Batten Disease

CLN6-Batten disease, a form of neuronal ceroid lipofuscinosis is a rare lysosomal storage disorder presenting with gradual declines in motor, visual, and cognitive abilities and early death by 12–15 years of age. We developed a self-complementary adeno-associated virus serotype 9 (scAAV9) vector expressing the human CLN6 gene under the control of a chicken β-actin (CB) hybrid promoter. Intrathecal delivery of scAAV9.CB.hCLN6 into the cerebrospinal fluid (CSF) of the lumbar spinal cord of 4-year-old non-human primates was safe, well tolerated, and led to efficient targeting throughout the brain and spinal cord. A single intracerebroventricular (i.c.v.) injection at postnatal day 1 in Cln6 mutant mice delivered scAAV9.CB.CLN6 directly into the CSF, and it prevented or drastically reduced all of the pathological hallmarks of Batten disease. Moreover, there were significant improvements in motor performance, learning and memory deficits, and survival in treated Cln6 mutant mice, extending survival from 15 months of age (untreated) to beyond 21 months of age (treated). Additionally, many parameters were similar to wild-type counterparts throughout the lifespan of the treated mice.

CLN6 is a 311-amino acid protein with seven predicted transmembrane domains, and it is predominately localized to the endoplasmic reticulum.4,5 As with other CLN proteins, its exact function remains unclear; however, it has been implicated in intracellular trafficking and lysosomal function.6–8 There are currently over 70 characterized disease-causing mutations in CLN6,13 with most of these mutations leading to either a complete loss of CLN6 protein or production of truncated CLN6 protein products that are thought to be highly unstable and/or non-functional.14 Several naturally occurring animal models of CLN6-Batten disease have been described,15 these include sheep,16,17 canine,18 and mouse models.19 The spontaneous mutation found in the Cln6exf mouse model, the model used in this study, recapitulates many of the pathological and behavioral aspects of the disease.20 The Cln6exf mice contain an insertion of an additional cysteine (c.307insC, frameshift after P102), resulting in a premature stop death of neurons contributes to the clinical signs and symptoms. Disease age of onset and disease progression vary considerably depending on the genetic variant of the disease. Here, we focus on CLN6-Batten disease, which can occur as two different forms: variant late-infantile (vLINCL), the more common form, and adult-onset NCL (also called type A Kufs disease).21 With vLINCL (referred to here as CLN6-Batten disease), age of onset is between 18 months and 6 years and death typically occurs by age 12–15. CLN6-Batten disease initially presents as impaired language and delayed motor and cognitive development in early childhood, with most patients being wheelchair bound within 4 years of disease onset.7 The disease progresses to include visual loss, severe motor deficits, recurrent seizures, dementia, and other neurodegenerative symptoms.

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hCLN6 Transcript and Protein Expressions Are Sustained through 18 Months of Age

To express human CLN6 (hCLN6), we designed an scAAV9 viral vector with the hCLN6 gene under control of the CB hybrid promoter, noted as scAAV9.CB.CLN6 (Figure S1A). This specific vector was chosen based on its CNS transduction efficiency and its proven safety in human subjects, as documented in a current phase I clinical trial for infants with type I spinal muscular atrophy.22 After cloning, transgene mRNA and protein expressions were verified in vitro in HEK293 cells (Figure S1B). Given the overarching concern in the Batten disease research community that overexpression of NCL proteins (including CLN6, CLN3, and CLN8) might be toxic to cells, we assessed this possibility by overexpressing hCLN6 in excitatory projection neurons of the cortical plate using a non-viral delivery method.

Cortical neurons were targeted using in utero i.c.v. electroporation of embryonic day 15.5 animals. Animals were electroporated with either scAAV9.CB.CLN6 or scAAV9.CB.GFP plasmids (1 µg/animal) and sacrificed at post-natal day 14 (P14), and the cellular trafficking of hCLN6 was examined. The additional GFP vector was used to visualize the distribution of scAAV9.CB.CLN6 in comparison to a non-NCL-expressing vector, indicating no abnormalities with hCLN6 distribution in cortical neurons in vivo (Figure S1C). Additionally, hCLN6 appeared to be properly trafficked in cortical neurons, and both vectors demonstrated robust expression in the cerebral cortex in vivo, indicating that overexpression of hCLN6 in cortical neurons alone does not appear to be detrimental to cells (Figure S1C).

The efficacy of CLN6 gene delivery following a single i.c.v. injection of scAAV9.CB.CLN6 (5 x 10^{10} vg/animal) into the CSF at P1 was determined in male and female Cln6nclf mice. Extensive details on animal usage and study design can be found in the Materials and Methods. These mice are a naturally occurring mouse model of CLN6-Batten disease, displaying major disease hallmarks found in human patients.19,20 Examination of hCLN6 expression by RT-PCR at 2, 6, and 18 months post-injection demonstrated sustained, robust hCLN6 expression in the cortex of scAAV9.CB.CLN6-injected Cln6nclf mice compared to PBS-injected controls (Figure 1A; Figure S2A), similar to previously reported scAAV9-CB-GFP expression levels.23-25 To examine the regional distribution of transgene expression, a modified in situ hybridization method called RNAscope was used to visualize hCLN6 transcript. scAAV9.CB.CLN6-injected mutant mice (i.c.v., P1, 5 x 10^{10} vg/animal) maintained a widespread expression of hCLN6 throughout many regions of the brain at 2, 6, and 18 months, including the somatosensory cortex and ventral postero medial (VPM)/ventral posterolateral (VPL) nuclei of the thalamus, two regions that have been shown to be affected earliest in the disease progression of the Cln6nclf mice (Figure 1B; Figures S2B and S3A).

To examine the expression of hCLN6 protein within the CNS, immunoblotting of cortical brain lysates harvested from scAAV9.CB.CLN6-injected Cln6nclf mice (i.c.v., P1, 5 x 10^{10} vg/animal) and PBS-injected controls was performed using anti-hCLN6 antibodies. Similar to what was seen with RNA expression, robust hCLN6 protein expression was seen throughout the CNS at 2, 6, and 18 months post-injection (Figure 1A; Figure S2A). Furthermore, immunolabeling of brain tissue using anti-hCLN6 antibodies confirmed the expression throughout the brain of scAAV9.CB.CLN6-treated Cln6nclf mice, particularly in the VPM/VPL and somatosensory cortex (Figure 1C; Figures S2B, S3B, and S4). Together, these findings demonstrate that CSF delivery of scAAV9.CB.CLN6 via i.c.v. injection is able to stably produce hCLN6 transcript and protein in disease-relevant regions of the CNS.

Sustained Expression of hCLN6 Prevents Classic CLN6-Batten Disease Pathology in the Brain

Although the pathogenicity of autofluorescent storage material (ASM) is not entirely understood, the accumulation of ASM is a hallmark of all forms of Batten disease. At 2, 6, and 18 months post-treatment, Cln6nclf mice injected with scAAV9.CB.CLN6 (i.c.v., P1, 5 x 10^{10} vg/animal) had a reduced accumulation of ASM within the VPM/VPL nuclei of the thalamus and somatosensory cortex of the brain compared to PBS-injected mice (Figure 2A; Figure S2C). Because PBS-treated Cln6nclf mice die by 15 months of age (as demonstrated in Figure 5D), moribund 12- to 14-month-old PBS-treated Cln6nclf mice were used as a comparison to 18-month-old scAAV9.CB.CLN6-treated Cln6nclf mice. Notably, the amount of ASM accumulation in these 18-month-old scAAV9.CB.CLN6-injected Cln6nclf mice was comparable to the age-matched untreated wild-type mice. One primary constituent of
the storage material is mitochondrial ATP synthase subunit C. At 2, 6, and 18 months of age post-injection, Cln6<sup>off</sup> mice treated with scAAV9.CB.CLN6 (i.c.v., P1, 5 × 10<sup>10</sup> vg/animal) had significantly reduced levels of ATP synthase subunit C accumulation within the VPM/VPL and somatosensory cortex of the brain, compared to control Cln6<sup>off</sup> mice injected with PBS (Figure 2B; Figure S2C).

Apart from the accumulation of storage material, other histological markers of disease progression include reactive gliosis, which occurs later in disease progression, and the reduction in dendritic spine density. At 6 and 18 months of age post-injection, Cln6<sup>off</sup> mice treated with scAAV9.CB.CLN6 (i.c.v., P1, 5 × 10<sup>10</sup> vg/animal) had significantly reduced astrocyte activation (glial fibrillary acidic protein [GFAP]) and microgliosis (CD68) in the VPM/VPL and somatosensory cortex as compared to moribund PBS-treated Cln6<sup>off</sup> mice (Figures 3A and 3B; the insets in Figure 3B show the morphology of microglia). To examine whether delivery of scAAV9.CB.CLN6 affected specific neuronal degeneration in Cln6<sup>off</sup> mutant mice, we examined dendritic spine density using classic Golgi-Cox impregnation techniques. Importantly, a single injection of scAAV9.CB.CLN6 (i.c.v., P1, 5 × 10<sup>10</sup> vg/animal) prevented early dendritic spine loss in 2-month-old Cln6<sup>off</sup> mice (Figure 4A).

As hCLN6 expression via AAV9 prevented many of the classic Batten disease pathologies at several time points and considering CLN6 is a transmembrane protein, we were interested in determining whether hCLN6 expression prevented these pathologies in only the cells that were transduced or if neighboring cells were also protected. When examining individual cell burden for storage material, we found that cortical cells from 18-month-old scAAV9.CB.CLN6-injected Cln6<sup>off</sup> mice (i.c.v., P1 injection, 5 × 10<sup>10</sup> vg/animal) were only protected from storage material burden if hCLN6 was present in that particular cell (Figure 4B). As such, while a single injection delivering scAAV9.CB.CLN6 into the CSF at P1 can prevent many of the classic CLN6-Batten disease pathologies in the brains of Cln6<sup>off</sup> mice, it appears that this effect is likely contained to the cells that are transduced upon delivery.

**Sustained Expression of hCLN6 Corrects Many of the Behavioral Deficits in Cln6<sup>off</sup> Mice**

Our previous work demonstrated that the Cln6<sup>off</sup> mouse model of Cln6-Batten disease recapitulates many of the motor, cognitive, and survival defects seen in humans. In the current study, using the rotarod as a classic measure of motor coordination, PBS-injected Cln6<sup>off</sup> mice began to show a decline in rotarod performance at 8 months of age compared to wild-type. However, the injection of Cln6<sup>off</sup> mice with scAAV9.CB.CLN6 (i.c.v., P1, 5 × 10<sup>10</sup> vg/animal) prevented this decline, an effect that lasted for the duration of the entire study period (24 months) (Figure 5A). To further study the effects of motor
coordination in detail, animals were subjected to various motor tasks (hind limb clasping, ability to lower oneself from a ledge, and gait assessment) at 12, 18, and 24 months of age, and they were assessed using a scoring matrix, with the highest score indicating the worst prognosis. Compared to PBS-treated Cln6nclf mice, mice treated with scAAV9.CB.CLN6 (i.c.v., P1, 5 × 10^10 vg/animal) showed significantly lower combined scores at all time points, with a slight increase in their score only at 24 months of age (Figure 5B).

As a measure of memory and learning abilities, mice were taken through a classic Morris water maze until 24 months of age. PBS-treated Cln6nclf mice performed poorly at the task starting at 9 months of age, indicated by their reduced ability to find the hidden platform, and treatment of Cln6nclf mice with scAAV9.CB.CLN6 (i.c.v., P1, 5 × 10^10 vg/animal) corrected this memory and learning deficit up to 12 months post-injection (Figure 5C). However, it is worth noting that the swim speeds of PBS-treated Cln6nclf mice were significantly reduced at 11 and 12 months of age, limiting the conclusions we could draw on memory and learning abilities of untreated animals at these later time points (Figure S5A). When comparing wild-type mice to scAAV9.CB.CLN6-treated animals (i.c.v., P1, 5 × 10^10 vg/animal) at later time points in the Morris water maze test, we found that even the treated mice needed more time to find the platform at 18 and 24 months, while the swim speed was the same among all test groups (Figure 5C; Figure S5A). To assess more subtle aspects of memory and learning, we also subjected mice to a water maze reversal test at 12, 18, and 24 months of age, where the platform was moved to a novel location. Cln6nclf mice treated with scAAV9.CB.CLN6 (i.c.v., P1, 5 × 10^10 vg/animal) took significantly longer to find the new platform location compared to wild-type mice in this test as well (Figure S5B). Taken together, these results indicate that a single treatment of scAAV9.CB.CLN6 prevents many of the motor declines seen in these animals, but it does not fully ward off memory and learning deficits when the mice are tested at later time points.

**Single i.c.v. Injection of scAAV9.CB.CLN6 Substantially Increases the Survival of Cln6nclf Mice**

We have previously shown that Cln6nclf mice have reduced survival compared to their wild-type counterparts. We compared the survival of scAAV9.CB.CLN6- and PBS-injected Cln6nclf mice with PBS-injected wild-type mice. A single i.c.v. injection of scAAV9.CB.CLN6 (5 × 10^10 vg/animal) into the CSF of Cln6nclf mice at P1 significantly increased animal survival compared to PBS-injected Cln6nclf mice (Figure 5D). While the median survival of PBS-treated animals was 14 months, scAAV9.CB.CLN6-treated mice had a median survival of 21.5 months. This is a highly significant 65% increase in survival rate. Moreover, the survival curve of scAAV9.CB.CLN6-treated Cln6nclf mice was not significantly different from wild-type animals.

Further, as a measure of overall health, body weight was recorded monthly. The improvement in health and survival was also underlined by the ability of scAAV9.CB.CLN6-treated mice to maintain their body weight, as no difference was observed compared to wild-type animals, while PBS-treated Cln6nclf mice started losing weight around month 12 (Figure S5C).
As a part of investigational new drug (IND)-enabling studies, we also performed a safety study with 172 wild-type mice treated with PBS and 223 wild-type mice treated with $5 \times 10^{10} \text{vg/animal}$ scAAV9.CB.CLN6 (i.c.v., P1), and we found that scAAV9.CB.CLN6 was well tolerated up to 24 weeks with no adverse effects attributable to the virus (data not shown). Taken together, we report the longest survival extension in the $Cln6nclf$ mouse model to date, and the data indicate the utility of a single treatment of scAAV9.CB.CLN6 to restore both cellular and functional deficits of CLN6-Batten disease.

Lumbar Intrathecal CSF Delivery of scAAV9.CB.CLN6 in Three 4-Year-Old Non-human Primates Is Safe and Well Tolerated

To test the safety of this treatment in a large animal model more relevant to human patients, we injected three 4-year-old male cynomolgus macaques with scAAV9.CB.CLN6. The animals were sacrificed at 1, 3, or 6 months post-injection. Each individual received a single lumbar intrathecal injection, delivering the viral vector directly into the CSF at a dose of $6 \times 10^{13}$ viral particles/animal. After the injection, the animals were held in a Trendelenburg position for 15 min, with head facing downward in a 45-degree angle to facilitate targeting of the brain and upper spinal cord areas, as previously demonstrated by our laboratory.25 All subjects recovered well from the injection and did not show any abnormal behavior.

A high expression of the transgene was found throughout the brain and spinal cord of all three animals (Figures 6A and 6B). Hematology and serum chemistry were performed at up to 5 time points during the study (baseline and 1, 2, 3, and 6 months), and they did not reveal major abnormalities. In particular, no evidence of elevation in aspartate aminotransferase (AST) or alkaline phosphatase (Alk Phos) enzyme levels was found, while alanine aminotransferase (ALT) was slightly increased in one animal at 1 month post-injection (below 200 U/L) (Figure 6C). No changes were found in total protein levels, creatinine, triglycerides, glucose, or ions such as phosphorus, calcium, magnesium, or sodium levels. Extensive histopathology was performed for each animal at the time of sacrifice. No abnormalities were found in any tissue analyzed, including various brain and spinal cord regions, heart, lung, liver, spleen, kidney, small intestine, skeletal muscles (diaphragm, triceps, tibialis anterior [TA], and gastrocnemius), and gonads, except for one animal that displayed a bladder infection at the time of necropsy. Together, our data indicate that the treatment with scAAV9.CB.CLN6 was well tolerated and safe in all three animals tested.

**DISCUSSION**

CLN6-Batten disease is a devastating neurodegenerative disease for which there is no cure or treatment available. CLN6 is ubiquitously expressed throughout the body, and it is unclear why neurons are more vulnerable to a loss of CLN6 compared to other cell populations.27 Identification of potential therapeutic targets has been hindered because the functions and mechanisms of action of CLN6 are largely unknown.28,29 Mutations in the CLN6 gene that result in CLN6-Batten disease are largely thought to be loss-of-function mutations caused by truncation of the protein and its subsequent rapid degradation.31,33 Reintroducing a functional copy of CLN6 using an AAV-mediated gene delivery is an attractive approach that does
not require precise knowledge of CLN6 function, and it has been explored in varying approaches in several NCLs.

The successful use of AAV-mediated gene therapy is dependent upon many factors, including careful selection of the AAV serotype and its tropism, the method of administration, the animal model and its age, the amount of virus, and the promoter driving gene expression. The promoters driving gene expression should be selected based on the desired cellular targets and the levels of expression desired. In the Bosch et al. study, they found that a low-expressing promoter, methyl-CpG-binding protein 2, driving Cln3 expression resulted in better outcomes compared to a high-expressing beta-actin promoter in a mouse model of CLN3-Batten Disease. The hybrid chicken beta-actin promoter was chosen for this study based on its ability to robustly drive gene expression. The promoters driving gene expression should be selected based on the desired cellular targets and the levels of expression desired. In the Bosch et al. study, they found that a low-expressing promoter, methyl-CpG-binding protein 2, driving Cln3 expression resulted in better outcomes compared to a high-expressing beta-actin promoter in a mouse model of CLN3-Batten Disease. The hybrid chicken beta-actin promoter was chosen for this study based on its ability to robustly drive gene expression in the CNS while being well tolerated.

There are many AAV serotypes, including recombinant AAV serotypes, which differ in their tropism (reviewed in Castle et al.). The AAV9 vector was chosen here for its ability to robustly target neurons and non-neuronal cells throughout the CNS. In addition to the tropism, the timing and method of injection can influence the biodistribution of the virus. In the mouse, at P1, the tropism of the AAV9 virus targets neurons and glia equally. After P1, the tropism of AAV9 in mice begins to shift from neurons and begins to preferentially target glia, a phenomenon that does not seem to occur in non-human primates.

Studies from the Kaspar lab and others have shown that both intravenous and i.c.v. administrations of AAV9 result in viral transduction throughout the CNS. However, when the Kaspar group and others used scAAV9.CBA.SMN to treat spinal muscular atrophy in SMNΔ7 mice, intravenous (3.3e14 vg/kg at P1) administration required ten times more virus than an i.c.v. (3.3e13 vg/kg) delivery to achieve rescue of the spinal muscular atrophy (SMA) phenotype. Using the minimum amount of virus to achieve efficacy in treatment of the disease is important, because several studies have shown that using too much virus can result in toxicity. Using the i.c.v. delivery, we were able to limit our viral injection to $5 \times 10^{10}$ vg/animal, and, by injecting directly into the CSF, more of the virus was retained in the CNS, reducing the exposure of organ systems outside the CNS to the virus itself. Our safety studies included 395 mice and three non-human primates, and we found that scAAV9.CB.CLN6 was well tolerated and safe in both mammalian model systems.

In this study, we demonstrate that a single i.c.v.-mediated injection of scAAV9.CB.CLN6 ($5 \times 10^{10}$ vg/animal) into the CSF of P1 mice was sufficient to induce stable, robust expression of hCLN6 protein throughout the CNS for up to 18 months. While it is tempting to question the clinical relevance of this approach, particularly the timing of treatment, we believe that the efficacy of this treatment will spur earlier diagnostic testing in the future (i.e., prenatal screenings), thereby making our early treatment protocol extremely relevant. Further, we anticipate the ongoing clinical trial to show slowed or halted disease progression in symptomatic patients treated with this vector, which may markedly improve their quality of life, even if not fully curative.

While we demonstrate robust mRNA and protein expressions of hCLN6, there are certainly inconsistencies across gene and protein expression, time points, and brain regions. Unfortunately, we cannot accurately compare RNA and protein expressions across time points, as each cohort of mice was immunolabeled in separate batches. However, we do believe that any differences between the time points are likely related to individual animal variability. Differences between mRNA and protein expressions are similarly hard to determine, though there are many possible explanations, including animal and injection variability, conditions of RNAscope incubation versus conditions of immunolabeling conditions, and the possibility of our RNAscope probes/hCLN6 antibody only partially detecting actual signal. As such, we cannot definitively conclude on the precise stability of hCLN6 expression, though this may be the subject of future study.

Additionally, the injection of scAAV9.CB.CLN6 in Cln6mut mice reduced many of the pathological hallmarks of CLN6-Batten disease and corrected many of the behavioral deficits characteristic of this...
model, with the exception of the Morris water maze test. Importantly, Batten disease also affects visual capabilities, which we did not explore in this particular publication. It is possible that visual deficits may have played a role in scAAV9.CB.CLN6-treated mice performing poorly in the water maze task, and, as such, the ability of a single, i.c.v. injection of scAAV9.CB.CLN6 to rescue visual deficits will remain the subject of future studies. In addition to improving these behavioral deficits, the administration of scAAV9.CB.CLN6 increased the longevity of the Cln6\textsuperscript{nclf} mice. None of the Cln6\textsuperscript{nclf} PBS-injected mice survived past 15 months, whereas a single injection of scAAV9.CB.CLN6 in the Cln6\textsuperscript{nclf} animals at P1 resulted in a 225-day increase in median survival, which to our knowledge is the largest increase in survival in a Batten disease mouse model using AAV-mediated gene therapy.

Taken together, these results indicate the outstanding efficacy of a single, i.c.v. injection of scAAV9.CB.CLN6 into the CSF of P1 Cln6\textsuperscript{nclf} mice and safety of scAAV9.CB.CLN6 in both mice and primates. Based on these promising data, this gene therapy construct has moved forward for expression and pathology assays as they became moribund, and 3 wild-type (WT) and 3 treated Cln6\textsuperscript{nclf} animals (mixed sexes) were sacrificed at 18 months of age to compare to the moribund Cln6\textsuperscript{nclf} tissues. Assays at this endpoint included qPCR, western, RNAscope, hCLN6 immunohistochemistry [IHC], Batten disease histopathology [ASM, SubC, GFAP, and CD68], and dendritic spine density). Cohort 2 consisted of 2–3 animals per sex per treatment group, and they were used for all 6-month expression and pathology assays (qPCR, western, RNAscope, hCLN6 IHC, and Batten disease histopathology [ASM, SubC, GFAP, and CD68]). Lastly, cohort 3 consisted of 10–14 animals per sex per treatment group; they were used for all behavioral assays from 3 to 24 months of age and monitored for survival. From this same cohort (cohort 3), 2 male and 2 female untreated Cln6\textsuperscript{nclf} animals were used for expression and pathology assays as they became moribund, and 3 wild-type (WT) and 3 treated Cln6\textsuperscript{nclf} animals (mixed sexes) were sacrificed at 18 months of age to compare to the moribund Cln6\textsuperscript{nclf} tissues. Assays at this endpoint included qPCR, western, RNAscope, hCLN6 IHC, and Batten disease histopathology (ASM, SubC, GFAP, and CD68). Taken together, assays from each time point were conducted on the same cohort of animals, and it was only the endpoint cohort that had any behavior testing before tissue was assessed for expression and pathology. Additionally, all assays were run independently on each cohort as samples became available, which limits any correlation between time points for the same assay.

**MATERIALS AND METHODS**

**Ethics Statement and Animals**

Wild-type and homozygous Cln6-mutant mice (Cln6\textsuperscript{nclf}) on C57BL/6J backgrounds were used for all studies and were housed under identical conditions. Mice received a single i.c.v. injection of either PBS or scAAV9.CB.CLN6 (5 × 10\textsuperscript{10} vg/animal) at P1 (the day after birth), following hypothermia sedation. Animals were monitored continuously until fully recovered from sedation and daily thereafter. All animals were genotyped using previously described techniques.\textsuperscript{20}

There were several separate cohorts of mice used throughout the study, all receiving a single i.c.v. injection of either PBS or scAAV9.CB.CLN6 (5 × 10\textsuperscript{10} vg/animal) at P1. Cohort 1 consisted of 4–5 animals per sex per treatment group, and they were used for all 2-month expression and pathology assays (qPCR, western, RNAscope, hCLN6 immunohistochemistry [IHC], Batten disease histopathology [ASM, SubC, GFAP, and CD68], and dendritic spine density). Cohort 2 consisted of 2–3 animals per sex per treatment group, and they were used for all 6-month expression and pathology assays (qPCR, western, RNAscope, hCLN6 IHC, and Batten disease histopathology [ASM, SubC, GFAP, and CD68]). Lastly, cohort 3 consisted of 10–14 animals per sex per treatment group; they were used for all behavioral assays from 3 to 24 months of age and monitored for survival. From this same cohort (cohort 3), 2 male and 2 female untreated Cln6\textsuperscript{nclf} animals were used for expression and pathology assays as they became moribund, and 3 wild-type (WT) and 3 treated Cln6\textsuperscript{nclf} animals (mixed sexes) were sacrificed at 18 months of age to compare to the moribund Cln6\textsuperscript{nclf} tissues. Assays at this endpoint included qPCR, western, RNAscope, hCLN6 IHC, and Batten disease histopathology (ASM, SubC, GFAP, and CD68). Taken together, assays from each time point were conducted on the same cohort of animals, and it was only the endpoint cohort that had any behavior testing before tissue was assessed for expression and pathology. Additionally, all assays were run independently on each cohort as samples became available, which limits any correlation between time points for the same assay.

All non-human primates were housed at the Mannheimer Foundation (Homestead, FL, USA), an assessment and accreditation of laboratory animal care (AAALAC)-accredited facility following the specifications recommended in The Guide for the Care and Use of Laboratory Animals (National Academy Press, Washington, DC, 1996). Each individual received a single lumbar intrathecal injection, delivering the viral vector directly into the CSF at a dose of 6 × 10\textsuperscript{13} viral particles per animal. After the injection, the animals were held in a Trendelenburg position for 15 min with head facing downward in a 45-degree angle to facilitate targeting of the brain and upper spinal cord areas, as previously demonstrated by our laboratory.\textsuperscript{27} All subjects recovered well from the injection and did not show any abnormal behavior. For...
platelet count and liver enzyme analysis, non-human primates were bled via the femoral vein, with non-terminal bleeds extracting no more that 1% body weight. Hematology and serum chemistry were analyzed by a third party company (Antech Diagnostics).

Vector
A human CLN6 cDNA clone was obtained from OriGene (Rockville, MD). hCLN6 cDNA was further subcloned into an AAV vector under the hybrid chicken β-actin promoter and tested

**Figure 6. scAAV9.CB.CLN6 Is Highly Expressed and Well Tolerated in Non-human Primates**

(A) A single lumbar intrathecal injection delivering scAAV9.CB.CLN6 into the cerebral spinal fluid (6 × 10^13 viral particles/animal) induces a high expression of the transgene throughout the brain and spinal cord of non-human primates, as shown by fluorescent western blot. Blots representative of 3 animals, with "+" indicating an animal with scAAV9.CB.CLN6 treatment. Ctx., cortex; C. Call., corpus callosum; P.V.W.M., periventricular white matter; Hipp., hippocampus; Cere., cerebellum; Thal., thalamus; Cervical, cervical spinal cord; Thoracic, thoracic spinal cord; Lumbar, lumbar spinal cord. (B) Quantification of fluorescent western blots in (A). Mean ± SEM; n = 3. Unpaired Student’s t test, *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001. (C) Delivery of scAAV9.CB.CLN6 (single lumbar intrathecal, 6 × 10^13 viral particles/animal) did not alter platelet concentration or elevate liver enzymes in the majority of non-human primates. Red data points indicate scAAV9.CB.CLN6-treated animals; blue data points indicate PBS-treated animals. ALT, alanine aminotransferase; AST, aspartate aminotransferase; Alk Phos, alkaline phosphatase; GGT, gamma-glutamyl transferase.
in vitro and in vivo. The scAAV9.CB.CLN6 was produced by transient transfection procedures using a double-stranded AAV2-inverted terminal repeat (ITR)-based CB-CLN6 vector, with a plasmid encoding the Rep2Cap9 sequence, as previously described, along with an adeno viral helper plasmid pHHelp (Stratagene, Santa Clara, CA) in HEK293 cells. The purity and titer of the vector were assessed by silver staining and qPCR analysis.

RT-PCR

Mice were CO2 euthanized and a lateral section of the cortical brain was collected and frozen for RNA isolation. Total RNA was extracted using a Maxwell 16 Mdx instrument and the Maxwell 16 LEV simplyRNA tissue kit (Promega), according to the manufacturer’s protocol. RNA quality and concentration were assayed using a BioTek Epoch Microplate Spectrophotometer, and only samples with concentrations between 200 and 1,000 ng/µL and an A260/A280 > 1.8 were used. Total RNA (1 µg) was used for reverse transcription using the Promega GoScript Reverse Transcription System (A5001) and following the manufacturer’s suggested protocol. PCR was performed using 0.5 µL cDNA and 400 nM primers (hCLN6 forward 5'-AAC GTC ATC ACG CCC TTT CT-3', hCLN6 reverse 5'-GAA GAG CAG CCG GTG GTT G-3', Gapdh forward 5'-ACC ACA GTC CAT GCC ATC AC-3', and Gapdh reverse 5'-ACC ACA GTC CAT GCC ATC AC-3') in a 25-µL reaction using a 3-step PCR reaction for 30 cycles and an annealing temp of 60°C. The hCLN6 RT-PCR probes had little cross-reactivity with mouse Cln6, as demonstrated in Figure 1A. PCR products were run on a 1% agarose gel and stained with ethidium bromide. Bands were visualized under UV light using a Gel-Doc Imager (UVP). Densitometry was performed with VisionWorks software (UVP).

In Utero Electroporation

All surgeries used sterile technique and were performed with approval from Sanford Research institutional animal care and use committee (IACUC). Briefly, dams at embryonic day (E)15.5 of pregnancy were anesthetized with 3% isoflurane, then a 1- to 2-cm abdominal midline incision was made. The uterine horns were gently pulled out to expose the pups and kept moist with sterile saline. Approximately 2 µL endotoxin-free, sterile plasmid solution was injected through the uterine tissue into a lateral cerebral ventricle of each pup using a 30G needle, followed by electroporation with five pulses of 50 V at 500-ms intervals via 3-mm diameter disc electrodes. The uterine horns were placed back into the dam, and the abdomen was remoistened. The muscle and skin layers were closed using a 6-0 silk suture with a simple interrupted pattern. Triple antibiotic ointment was applied to the incision site, and the dam was placed in a pre-warmed cage to recover from anesthesia.

CLN6 Transcript Detection Using RNAscope

Mice were CO2 euthanized and cardiac perfused with PBS. Brains were collected and placed on a 1-mm sagittal brain block. Brains were sliced at the midline and 3 mm right of the midline. The 3-mm sagittal piece was flash frozen with −50°C isopentane and then sectioned on a cryostat at 16 µm and placed on slides. Slides were then processed according to the manufacturer’s suggested protocols (ACDBio manuals 320293 and 320513). Sections were labeled with a human-specific CLN6 probe (ACDBio 452478), which consisted of 6 double Z pairs in regions of the Cln6 gene with little homology between mouse and human CLN6. The hCLN6 probes had little to no cross-reactivity with mouse Cln6, as demonstrated in Figure 1B (wild-type panels). Slides were fluorescently labeled with the RNA-scope Fluorescent Multiplex Kit (ACDBio 320850) using their Amp 4-FL-A1B, which tagged the hCLN6 probe with a 550-nm fluorophore; slides were counterstained with DAPI to label nuclei. Tissue sections were mounted on slides under coverslips using antifade mounting media (Dako faramount, Agilent). Slides were stored in the dark before imaging.

Immunoblotting and Histopathology

Wild-type and Cln6−/− mice were CO2 euthanized, perfused with PBS, and tissue was either frozen or fixed with 4% paraformaldehyde (PFA). Immunoblotting was performed as previously described using anti-hCLN6 and anti-β-actin antibodies. Rabbit polyclonal anti-hCLN6 antibody was raised against a synthetic peptide corresponding to the N-terminal amino acids 1–20 of human CLN6. The hCLN6 antibodies had little to no cross-reactivity with mouse Cln6, as demonstrated in Figure 1C. Fixed brains were sectioned on a vibratome at 50 µm (Leica VT1000S). Sections were processed with standard immunofluorescence and 3,3′-diaminobenzidine (DAB)-staining protocols. The primary antibodies included anti-CD68 (AbD Serotec, MCA1957; 1:250), anti-GFAP (Dako, Z0334; 1:250), and anti-ATP synthase subunit C (Abcam, ab181243; 1:500). The secondary antibodies included anti-rat and anti-rabbit biotinylated (Vector Labs, BA-9400; 1:2,000) and Alexa Fluor fluorescent secondarys (1:1,500).

For the hCLN6 immunolabeling, 16-µm fixed sagittal brain sections were cut on a cryostat and processed with standard immunofluorescence staining protocols. Antibodies included anti-hCLN6 (1:250) and Alexa Fluor fluorescent secondaries (1:1,500). Sections were scored positive for the accumulation of storage material when more than three fluorescent puncta were aggregated around the nucleus. Subunit C, GFAP, and CD68 immunoreactivity was quantified using a threshold analysis in NIS-Elements Advanced Research software (version v.4.20). Images were taken in the VPM/VPL of the thalamus and layers 2 and 3 of the somatosensory cortex, with multiple images taken of multiple tissues from each animal. For autofluorescent storage material, cells were scored positive for the accumulation of storage material when more than three autofluorescent puncta were aggregated around the nucleus. Subunit C, GFAP, and CD68 immunoreactivity was quantified using a threshold analysis in NIS-Elements Advanced Research software (v.4.20).

For determining dendritic spine density, brains were processed using a standard Golgi-Cox protocol. First, tissues were incubated in 1% HgCl2, 1% K2Cr2O7, and 0.8% K2CrO4 solution for 14 days. Brains were then put through a 5-day sucrose gradient, sectioned at 50 µm on a vibratome, and incubated in 50% ammonia for 30 min. Finally, brain sections were washed and incubated in 1% Na2S2O3, washed,
mounted, and dehydrated through an ethanol and CitroSolv gradient prior to mounting on glass slides; 15 pyramidal neurons/animal were imaged from layers 5 and 6 of the cortex. Using NIS-Elements software, the length and number of dendritic spines of primary dendrites were measured using the following criteria: dendrites must be >25 μm, and spine counts must start within 35 μm of the cell body and terminate 225 μm from the cell body.

For non-human primate experiments, animals were CO₂ euthanized, perfused with PBS, and tissue was either frozen or fixed with 4% PFA. Immunoblotting was performed as previously described using anti-hCLN6 (1/500) and anti-GAPDH antibodies (Millipore, 1/5,000). Briefly, tissue was lysed using tissue protein extraction reagent (TPER) buffer (Thermo Scientific) and complete protease inhibitor cocktail (Roche). 70 μg protein was separated by SDS-PAGE (NuPage Bis-Tris gels, Life Technologies) and transferred onto polyvinylidene fluoride (PVDF) membrane (Millipore). Membrane was blocked using Odyssey Blocking buffer (LI-COR Biosciences) and incubated with primary antibodies overnight. The following day the membrane was washed and incubated with LI-COR secondary antibodies (1/25,000), according to the manufacturer’s protocols, and imaged using Odyssey CLx (LI-COR Biosciences). Rabbit polyclonal anti-hCLN6 antibody was raised against a synthetic peptide corresponding to the N-terminal amino acids 1–20 of human CLN6.

Neurobehavior Testing

Rotarod
Study groups were tested monthly (at months 3–12, 18, and 24) on a Rotarod-5 Rotarod (Columbus Instruments, Columbus, OH, USA) to assess motor abilities. The machine was set to accelerate 0.3 rpm every 2 s, with a starting speed of 0.3 rpm and a maximum speed of 36 rpm. Mice were trained for three consecutive trials, given a 30-min rest period, trained for three consecutive trials, given a second 30-min rest period, and trained for three final consecutive trials. After a 4-h rest period, mice were tested using the same paradigm as the training session. The latency time to fall from the rod was averaged from each of the nine afternoon testing sessions to produce one value per mouse.

Water Maze
Mice were tested monthly (at months 3–12, 18, and 24) in a Morris water maze apparatus to assess memory and learning deficiencies. The apparatus consisted of a 4-ft-diameter tube filled with water to about 26 in, with the goal platform submerged by 0.5 cm. The tube was aligned with four distinct visual cues at 0, 90, 180, and 270 degrees, with the platform resting in the maze at 315 degrees. Mice were first trained in a clear pool with a flagged platform. Mice were given 60 s to complete each trial, with four trials in the morning, followed by a 3-h rest period, and four additional trials in the afternoon. Mice that could not locate the platform with 50% accuracy in the time allotted were eliminated from further testing. Mice were then tested in water colored with white, non-toxic tempura paint and an unflagged platform. Mice were given 60 s to complete each trial, with four trials in the morning, followed by a 3-h rest period, followed by four additional trials in the afternoon. Mice were tested for 4 consecutive days, each day starting at a different visual cue. Mice were recorded using Any-maze video tracking software (Stoelting, Wood Dale, IL, USA); test duration and swim speed are represented as the average from 16 afternoon trials per mouse. At 12, 18, and 24 months of age, an additional 4 days of reversal testing was introduced, where the hidden platform was moved from 315 to 45 degrees.

Clasping, Ledge, and Gait Tests
Tests were performed as previously described on 12-, 18-, and 24-month-old wild-type and Cln6nclf mice. Briefly, for hind limb-clasping measurements, animals were scored on the extent to which their limbs clasped into their abdomen when held by the base of their tail (score 0–3). For ledge-lowering measurements, animals were scored on their ability to easily lower themselves from the edge of their home cage (score 0–3). For gait measurements, animals were scored on their overall ease of walking, including whether their abdomen dragged on the ground and if their limbs were splayed while walking (score 0–3).

Statistical Analysis
Statistical analyses were performed using GraphPad Prism (v.6.04) and details are noted in the figure legends. In general, one-way ANOVA was employed with a Bonferroni correction, and outliers were removed with the ROUT method (Q = 0.1%). If appropriate, an unpaired t test was used. For the survival curve analysis, the log rank (Mantel-Cox) test was used. Statistical significance was indicated as follows: *p < 0.05, **p < 0.01, ***p < 0.001, and ****p < 0.0001. The datasets generated during and/or analyzed during the current study are available from the corresponding author upon reasonable request.

Study Approval
All mouse studies were performed in an AAALAC-accredited facility in strict accordance with NIH guidelines, and they were approved by the Sanford Institutional Animal Care and Use Committee (U.S. Department of Agriculture [USDA] license 46-R-0009).

SUPPLEMENTAL INFORMATION
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AUTHOR CONTRIBUTIONS
Conceived and Designed the Experiments, J.T.C., S.L., K.A.W., D.J.T., F.R., B.K.K., K.M., and J.M.W.; Provided Reagents, S.M.H. and S.Y.L.; Executed the Experiments, J.T.C., S.L., K.A.W., D.J.T., S.S.D., T.B.J., C.N.D.-R., F.R., D.M., S.C., P.M., C.P., and K.M.; Analyzed the Data, J.T.C., S.L., K.A.W., D.J.T., T.B.J., C.N.D.-R., F.R., D.M., S.C., P.M., C.P., K.M., and J.M.W.; Contributed to the Writing of the Manuscript, J.T.C., K.A.W., D.J.T., K.M., and J.M.W.; Agree with the Main Conclusions and Interpretations, J.T.C., S.L., K.A.W., D.J.T., S.S.D., T.B.J., C.N.D.-R., F.R., D.M., S.C., P.M., C.P., S.M.H., S.Y.L., B.K.K., K.M., and J.M.W.; Jointly Developed the Structure and Arguments for the Paper, J.T.C., K.A.W., D.J.T., K.M., B.K.K., and J.M.W.;
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