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Jacqueline A. Hawkins-Salsbury
Washington University School of Medicine in St. Louis

Lauren Shea
Washington University School of Medicine in St. Louis

Xuntian Jiang
Washington University School of Medicine in St. Louis

Daniel A. Hunter
Washington University School of Medicine in St. Louis

A Miguel Guzman
St. Louis University

See next page for additional authors

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Mechanism-Based Combination Treatment Dramatically Increases Therapeutic Efficacy in Murine Globoid Cell Leukodystrophy

Jacqueline A. Hawkins-Salsbury,1 Lauren Shea,1 Xuntian Jiang,2 Daniel A. Hunter,2 A. Miguel Guzman,4 Adarsh S. Reddy,1 Elizabeth Y. Qin,1 Yedda Li,1 Steven J. Gray,2 Daniel S. Ory,1 and Mark S. Sands1,3

Departments of 1Internal Medicine, 2Surgery, and 3Genetics, Washington University School of Medicine, St Louis, Missouri 63110, 4Department of Pathology, St Louis University School of Medicine, St Louis, Missouri 63104, and 3Department of Ophthalmology, University of North Carolina School of Medicine, Chapel Hill, North Carolina 27599

Globoid cell leukodystrophy (GLD, Krabbe disease) is a lysosomal storage disease (LSD) caused by a deficiency in galactocerebrosidase (GALC) activity. In the absence of GALC activity, the cytotoxic lipid, galactosylsphingosine (psychosine), accumulates in the CNS and peripheral nervous system. Oligodendrocytes and Schwann cells are particularly sensitive to psychosine, thus leading to a demyelinating phenotype. Although hematopoietic stem-cell transplantation provides modest benefit in both presymptomatic children and the murine model (Twitcher), there is no cure for GLD. In addition, GLD has been relatively refractory to virtually every experimental therapy attempted. Here, Twitcher mice were simultaneously treated with CNS-directed gene therapy, substrate reduction therapy, and bone marrow transplantation to target the primary pathogenic mechanism (GALC deficiency) and two secondary consequences of GALC deficiency (psychosine accumulation and neuroinflammation). Simultaneously treating multiple pathogenic targets resulted in an unprecedented increase in life span with improved motor function, persistent GALC expression, nearly normal psychosine levels, and decreased neuroinflammation. Treating the primary pathogenic mechanism and secondary targets will likely improve therapeutic efficacy for other LSDs with complex pathological and clinical presentations.

Key words: dysmyelination; gene therapy; globoid cell leukodystrophy; Krabbe disease; lysosomal storage disease

Introduction

Globoid cell leukodystrophy (GLD, Krabbe Disease) is an inborn error of metabolism resulting from a deficiency in the lysosomal enzyme galactocerebrosidase (GALC; Wenger et al., 2001). Loss of GALC activity results in the accumulation of a cytotoxic lipid, galactosylsphingosine (psychosine), in the CNS and peripheral nervous system leading to a dysmyelinating phenotype (Miyatake and Suzuki, 1973). The currently available treatment for GLD is hematopoietic stem-cell transplantation (HSCT), which provides only modest improvements if initiated before symptom onset (Krivit et al., 1998; Escolar et al., 2005; Martin et al., 2006). Therefore, the development of new and more efficacious therapeutic regimens is of the utmost importance.

Received Oct. 10, 2014; revised Feb. 23, 2015; accepted March 15, 2015.

An authentic mouse model of GLD, the Twitcher mouse, has been available for more than three decades (Kobayashi et al., 1980). This mouse mimics the human disease and has a rapid and severe disease course (Suzuki and Suzuki, 1983). Despite the availability of the Twitcher mouse, development of effective therapies has been slow. Virtually all single-therapy approaches, including BMT (Yeager et al., 1984), gene therapy (Lin et al., 2005, 2007; Galbiati et al., 2009; Gentner et al., 2010; Reddy et al., 2011; Rafi et al., 2012), substrate reduction (LeVine et al., 2000; Biswas and LeVine, 2002), antioxidants (Hawkins-Salsbury et al., 2012a), steroids (Kagitan-Shimono et al., 2005; Luzi et al., 2009), enzyme replacement (Lee et al., 2005; Qin et al., 2012), and neuronal (Pellegatta et al., 2006) or mesenchymal (Ripoll et al., 2011) progenitor cells, have met with minimal to modest success. GLD is a complex disease, involving the primary pathogenic mechanism and secondary consequences. Therefore, it is not surprising that single therapeutic approaches provide limited efficacy. Combination therapies that target both primary and secondary mechanisms have been more successful. Intracranial gene therapy targeting the primary defect synergized with bone marrow transplantation (BMT) to nearly triple the lifespan of the Twitcher mouse (Lin et al., 2007). Subsequent improvements in the route of gene therapy delivery further extended the life span (Reddy et al., 2011). Substrate reduction therapy for LSDs decreases the rate of accumulation of undegraded metabolites by reducing the synthe-
sis of the enzyme substrates. This can be accomplished in GLD with L-cycloserine, which broadly decreases the synthesis of sphingolipids, including psychosine (Sundaram and Lev, 1985; LeVine et al., 2000). L-cycloserine alone significantly improves lifespan in the Twitcher mouse to ~57 d (LeVine et al., 2000) and synergizes with BMT to increase lifespan to ~112 d (Biswas and LeVine, 2002).

Here we show that targeting the primary pathogenic mechanism and two secondary consequences of GALC deficiency using diverse therapeutic approaches dramatically increases treatment efficacy in the Twitcher mouse. Mice were treated from an early age with “triple therapy” consisting of BMT, adeno-associated virus 2/5 (AAV2/5)-mediated gene therapy, and substrate reduction with L-cycloserine. Interestingly, each approach increased efficacy in a synergistic manner, resulting in an unprecedented increase in lifespan and persistent clinical/behavioral improvements. Simultaneously targeting distinct pathogenic pathways may be the most effective way to treat complex inherited metabolic disorders, such as lysosomal storage diseases.

Materials and Methods

Experimental design. Treated and untreated Twitcher mice, as well as untreated controls were used in this study. Twitcher mice were treated with CNS-directed gene therapy on postnatal day (P)2, BMT on P3, and L-cycloserine three times a week starting on P5. Experimental and control animals of both sexes were randomly assigned to groups for analysis at predetermined time points (n = 3–6 animals/time point) or for life span and behavioral analyses (n = 10–14 animals/group). The number of animals/group was based on previous experience with this model and the specific assays, as well as power calculations. Quantitative biochemical analyses (GALC activity, psychosine measurements, cytokine/chemokine levels) were performed in duplicate on tissues from three to six animals, depending on the assay. Semiquantitative histomorphometry was performed on at least three representative sections/animal from at least three animals/group. The photography settings were kept constant for all images. ANOVA was used for group comparisons with Bonferroni correction for pairwise comparisons. Student’s t-test was used to compare the means of two groups. Significant differences in life span were determined by log-rank test. Differences were deemed significance at a p value of <0.05.

Experimental animals. Animals were housed at Washington University School of Medicine under the supervision of MSS. The Institutional Animal Care and Use Committee at Washington University School of Medicine approved all animal protocols. Heterozygous Twitcher (GALC +/−) mice on a C57BL/6 background were obtained from the Jackson Laboratory, and heterozygote by heterozygote matings were used to generate the homozygous Twitcher (GALC −/−) mice used in this study. Genotypes were determined on P1 by PCR as previously described (Sakai et al., 1996; Lin et al., 2005). Hematopoietic stem cell donors were sex-matched syngenic GALC +/+ mice expressing GFP under control of the long untranslated region from the rabbit β-globin gene. The single-stranded AAV2 vector was pseudotyped with an A500 capsid. Virus was produced using a triple-transfection method previously described. Briefly, one brain hemisphere or two sciatic nerves from a single mouse were homogenized in 0.1 M sodium citrate. Fifty microliter of each sample was added to 20 µl of N,N-dimethylpsychosine (250 ng/ml) internal standard and 200 µl MeOH. Samples were vortexed and centrifuged, and then the supernatant collected. This extraction was repeated on the remaining pellet and the supernatants pooled. Psychosine concentrations were obtained using a column-switching LC-MS/MS method. Detection was achieved using an AB SCIEX 4000QTRAP tandem mass spectrometer (Applied Biosystems/MDS Sciex) using ESI in the positive ion mode along with multiple reaction monitoring. Analyst software (v1.5.1, Applied Biosystems/MDS Sciex) was used for the data analysis. The calibration curves (analyte peak area/internal standard peak area for y-axis and analyte concentration for x-axis) of psychosine were obtained using the least square linear regression fit (y = ax + b) and a weighting factor of 1/x². The coefficient of determination (r²) was set as >0.98 for acceptance criteria of calibration curves.

GALC activity. Following perfusion, one brain hemisphere was homogenized in ddH2O. The brain homogenate was centrifuged and the supernatant collected. GALC activity was measured as previously described using a 3H-glucosylceramide (Lin et al., 2005). After the reaction, uncleaved substrate was removed through galactose saturated chloroform/methanol extraction. The remaining radioactivity, in the form of free 3H-galactose, was measured in a scintillation counter. The specific activity of GALC enzyme was calculated as nanomoles of substrate cleaved per hour per milligram of protein.

Lifespan, body weight, and behavioral testing. Lifespan was recorded as the age of the animal on the date of death or kill. Animals were killed when they became moribund as defined as one or more of the following: losing >25% of their maximal body weight, ataxia severe enough to impair ability to eat or drink, or lack of response to tactile stimulus. Unless otherwise noted, all animals were killed by anesthetic overdose and perfused transcardially with PBS until the liver was cleared of blood.

Bodyweight was recorded for each mouse (n = 12) once a week. Beginning at week 3, mice were treated every other week for performance on the rotarod and inverted wire-hang tests as previously described (Reddy et al., 2011). The maximum length of either test was 60 s. Each animal was given at least 10 min to recover between trials or tests. Rotarod: mice were gently placed on a stationary rod, and then the rotation started. The rod was set to rotate at a constant 3RPM. Latency to fall was recorded. The average of three trials was recorded. Wirehang: animals were placed up-
right on a rigid 1/4 inch wire mesh screen. The screen was gently shaken to encourage gripping then turned upside down 12 inches over soft bedding. Latency to fall was recorded.

**LFB/PAS staining and immunohistochemistry.** Brains were harvested immediately following perfusion and either flash frozen for biochemical analyses or fixed in 4% paraformaldehyde for 24 h at 4°C. The fixed brains were then moved to 30% sucrose in tris-buffered saline for an additional 24 – 48 h at 4°C. Following cryoprotection, the brain was embedded in optimal cutting medium (Sakura Finetek), frozen on dry ice, and cryosectioned. The 16 μm sections were stained with luxol fast blue and periodic acid Schiff (LFB/PAS) as previously described (Lin et al., 2005). Additional sections were immunostained with anti-GFAP, anti-CD68, or anti-GFP antibodies according to standard methods. Briefly, free-floating sections were incubated in 1% hydrogen peroxide to quench endogenous peroxidase activity. Sections were then blocked in normal goat serum and incubated in primary antibody. Sections were next incubated in the appropriate biotinylated secondary antibody, followed by incubation in Vectastain Elite ABC reagent (Vector Laboratories). Sections were developed using a DAB peroxidase substrate kit (Vector Laboratories). Antibodies were as follows: rabbit anti-mouse glial fibrillary acidic protein (GFAP; 1:100; Immunostar), rat-anti-mouse CD68 (1:1000; AbD Serotec), rabbit polyclonal anti-mouse GFP (1:10,000; Abcam), biotinylated anti-rabbit IgG supplied with the Vectastain kit (1:200; PK-6101 Vector Laboratories), and mouse-adsorbed biotinylated anti-rat IgG (1:200; Vector Laboratories). Stained sections were mounted, dehydrated and coverslipped. Images were captured using an Olympus BX41 microscope and Olympus DP20 camera. All image capture variables were kept identical. At least three sections per animal and three animals per treatment group were analyzed for each brain area studied. Staining was quantified using ImageJ software. The percentage of total image area with staining above a set threshold value was calculated as the area fraction.

**Sciatic nerve.** Sciatic nerves were removed and fixed in phosphate-buffered 4% paraformaldehyde/2% glutaraldehyde (n = 4 – 6 per treatment group). Nerves were then incubated in osmium tetroxide, serially dehydrated in ethanol, and embedded in Araldite 502 (Polysciences). An ultramicrotome was used to prepare 1 μm cross-sections, which were then stained with 1% toluidine blue dye and mounted on slides. A Hitachi CCD KP-M1AN digitizing camera mounted on a Leitz Laborlux S microscope was used for image acquisition. Histomorphometric analysis was performed using the Leco IA32 Image Analysis System as previously described (Hunter et al., 2007).

**Cytokines.** PBS-perfused brains (n = 4 – 10 per treatment group) were flash-frozen in liquid nitrogen, then thawed on ice and homogenized in a
Figure 2. Triple therapy supplies persistent GALC activity and decreases psychosine accumulation in the Twitcher mouse. A, GALC activity was increased ~3.5-fold from WT levels in triple-treated Twitcher mice at 36 d (n = 4–6 animals per group). Though a slight decrease in GALC activity was observed as the animals aged, GALC activity remained approximately twice WT levels in terminal triple-treated animals. B, Thrice weekly treatment with 25 mg/kg L-cycloserine alone resulted in a significant reduction in brain psychosine in Twitcher mice (n = 4–5 animals per group). C, Triple therapy significantly reduced whole-brain psychosine levels in Twitcher mice at 36 d. Psychosine levels remained low at terminal time points (n = 4–5 animals per group). D, Sciatic nerve psychosine levels in Twitcher mice were significantly reduced by triple therapy at every time point (n = 4–7 animals per group; **p < 0.01, ***p < 0.001).

Figure 3. Persistent donor chimerism is observed in the bone marrow and brains of triple-treated Twitcher mice. Rare GFP-positive donor-derived cells were present in the brainstem, cerebellum, and thalamus of 36 d triple-treated mice (36D Triple). In terminal triple-treated animals (Terminal Triple) larger numbers of GFP-positive cells were observed in brainstem, cerebellum, and thalamus, among other brain regions. There is a significant increase in donor bone marrow engraftment (right) in terminal triple-treated Twitcher mice (22.4%) compared with 36 d triple-treated Twitcher mice (9.9%) as determined by flow cytometry. (*p < 0.05).
solution containing 50 mM Tris-HCl, 2 mM PMSF, 2 mM EDTA, and protease inhibitors (P8340, Sigma-Aldrich) with an Eperbach Con-
Torque tissue homogenizer. Samples were centrifuged at 14,000 g for 4 min at 4°C and supernatant was sent for rodent multianalyte profile analysis (Myriad Rules-Based Medicine).

Statistical analysis. Statistical significance was calculated using a one-
way ANOVA followed by Bonferroni correction for multiple compari-
sions to compare all treatment groups unless otherwise specified.

Results

Lifespan

Untreated Twitcher animals had a median lifespan of 39.5 d (Fig. 1 top left). We previously showed that BMT or CNS-directed AAV-mediated gene therapy initiated in newborn Twitcher mice increased the median life span to ~45 and ~70 d, respectively (Lin et al., 2007; Reddy et al., 2011). In the current study, the median lifespan for mice treated with L-cycloserine alone or L-cycloserine plus CNS-directed gene therapy is ~57 and ~70 d, respectively. In contrast, triple-treated Twitcher mice had a median lifespan of 298.5 d, with a range of 160 – 454 d. Additionally, several of the treated male mice were able to impregnate normal females, whereas untreated Twitcher males are not able to breed. Treated females were housed exclusively with other females, as the effect of carrying a pregnancy was uncertain and not a goal of this study. Therefore, it remains unknown whether triple-treated Twitcher females might be fertile.

Body weight

Twitcher mice have impaired weight gain, with weight at death approximately one-half that of age-matched GALC +/- litter-
mates (an average of 8 vs 16 g; Fig. 1 top right). Triple-treated mice have increased weight gain compared with untreated Twitcher mice, though not to the same extent as WT animals (maximum weight ~18 vs 25 g for a WT mouse). The weight of triple-treated animals was stable until immediately preceding death.

Motor function

Twitcher mice have impaired motor function as assessed through the rotarod and inverted wire-hang tests. Untreated Twitcher mice are unable to perform a test of coordination (rotarod) by 3–4 weeks of age. On average, triple-treated mice can perform the rotarod test normally until ~30 weeks of age, then slowly decline in their ability (Fig. 1 bottom left). Although the mean latency to fall steadily decreases over time, individual triple-treated mice typically perform normally (60 s) until 1.5–2 months before they die. For example, two triple-treated Twitcher mice that lived to 450 and 454 d of age were able to stay on the rotarod for 60 s until 400 and 395 d, respectively. The inverted wire-hang is a test of grip strength and is used as a surrogate for peripheral nerve function. Untreated Twitcher mice are unable to hang onto an inverted wire screen for the full 60 s test even at an early stage of disease. Triple-treated animals can perform this task better than untreated mice, but still show significant deficits compared with WT animals (Fig. 1 bottom right).

GALC activity

Twitcher mice have virtually undetectable whole-brain GALC activity as measured by cleavage of the radio-labeled natural sub-
strate of GALC, galactosylceramide (Fig. 2A). At 36 d, the level of whole-brain GALC activity in triple-treated Twitcher mice is ~3.5-fold greater than WT animals. Although there is an apparent decrease in GALC activity as the animals age, this is not statistically significant. Even at a terminal time point, triple-treated Twitcher mice still have nearly double the WT levels of brain GALC activity.

Psychosine levels

Whole-brain psychosine levels are elevated in Twitcher mice relative to WT mice (Fig. 2B,C). Thrice-weekly injections of L-cycloserine alone significantly reduces whole-brain psychosine (Fig. 2B). Previous studies have shown that AAV2/5-mediated gene therapy alone reduces whole-brain psychosine levels, whereas BMT alone does not (Reddy et al., 2011). As expected, triple-treated Twitcher mice also show reduced psychosine levels in the brain compared with untreated Twitcher mice (Fig. 2C). Although whole brain psychosine levels in triple-treated animals are likely higher than WT, this difference is not statistically significant. Whole-brain psychosine levels remain low throughout the lifespan of triple-treated mice, with levels indistinguishable from WT at 160 d and at terminal time points. Psychosine also accumulates in the sciatic nerve of Twitcher mice. This accumulation is not observed in triple-treated Twitcher mice, even at terminal time points (Fig. 2D).

GFP-positive donor cell engraftment

By 36 d, triple-treated animals have an average of 9.9% (range = 5–30%) donor chimerism in the bone marrow (Fig. 3, Bone Marrow Engraftment). There is a significant increase in average donor bone marrow engraftment (22.4%) in terminal triple-treated mice. Engraftment of donor microglia in the brain is a much slower process (Kennedy and Abkowitz, 1997). At 36 d, only rare GFP-positive cells are observed in the brains of triple-treated mice. In contrast, numerous GFP-positive cells can be seen in...
terminal animals, particularly in the brainstem, cerebellum, and thalamus (Fig. 3).

Myelination and “globoid cell” infiltration
There is a substantial decrease in LFB staining in the cortex, brainstem, and cerebellum of untreated 36-d-old Twitcher mice compared with age-matched normal controls (Fig. 4). There is also an increase in the number of PAS-positive globoid cells in the untreated 36-d-old Twitcher mice compared with the normal controls. The level of LFB staining in 36-d-old triple-treated Twitcher mice is comparable to the normal control. Although there are still some globoid cells present in the 36-d-old triple-treated Twitcher mice it is greatly reduced compared with age-matched Twitcher mice. The intensity of LFB staining in 160-d-old triple-treated Twitcher mice is comparable to the 36-d-old untreated Twitcher mice. Although the number of PAS-positive cells in 160-d-old triple-treated Twitcher mice is comparable with 36-d-old untreated Twitcher mice, the cells appear smaller with less PAS-positive material.

Microglial and astrocyte activation
As previously reported, an increased number of CD68-positive macrophages and activated microglia are found in the brains of Twitcher mice compared with WT controls at 36 d (Lin et al., 2007; Reddy et al., 2011). Triple treatment dramatically decreased CD68 staining at 36 d (Fig. 5A). However, CD68 staining steadily increases as the treated mice age. A similar trend is observed for GFAP-positive activated astrocytes (Fig. 5B). Astrocytosis is diminished in triple-treated Twitcher brains at 36 d, but increases as the animals age.

Cytokine/chemokine levels
To further characterize the effect of triple therapy on neuroinflammation in the Twitcher brain, cytokine/chemokine levels were measured in whole-brain homogenates. Several proinflammatory cytokines and chemokines were increased in untreated Twitcher brains relative to WT at 36 d (n = 4–10 animals per group). Triple therapy resulted in a significant decrease in proinflammatory cytokines/chemokines. The levels remained indistinguishable from WT at terminal time points, with the exception of a slight increase in FGF-basic. IL-10, a cytokine with anti-inflammatory functions, was significantly increased in untreated Twitcher brains relative to WT. Triple-treated Twitcher brains showed IL-10 levels that were indistinguishable from WT at 36 d and at terminal time points (*p < 0.05, ****p < 0.0001).

Peripheral neuropathy
Untreated Twitcher mice demonstrate severe hindlimb weakness, such that hind limbs are essentially paralyzed when mice reach a terminal time point at ~40 d of age. Untreated Twitcher sciatic nerves are grossly swollen and display several features that are characteristic of Krabbe disease, including myelin loss, increased interstitial edema, cellular infiltration, and decreased axon density (Fig. 7). Quantitative histomorphometric analyses showed a significant decrease in axon density and a trend toward decreased myelin width in untreated Twitcher mice (Fig. 7). These changes are not prevented by triple therapy, and sciatic nerve pathology gets significantly worse over the lifespan of treated mice. Sciatic nerves from terminal triple-treated mice also show a unique histological pattern consisting of concentric swirls of myelin and connective tissue reminiscent of “onion bulb” formations. This pattern was not observed in 36 d triple-treated animals (Fig. 7).

Discussion
GLD has proven refractory to most therapeutic interventions. This is likely due to the complex nature of this disease, involving the primary and multiple secondary pathogenic mechanisms. In the current study, we targeted the primary defect in Twitcher...
mice with AAV-mediated, CNS-directed gene therapy (Fig. 8). In addition, we targeted two secondary consequences of GALC deficiency, psychosine accumulation and neuroinflammation, with substrate reduction therapy and BMT, respectively (Fig. 8). This multipronged approach resulted in the greatest increase in lifespan to date, with triple-treated Twitcher mice living to a median of ∼300 d and several mice living >450 d. Based on the increases in lifespan when applied singly, the combination of these three treatments should result in a median lifespan of ∼90 d if they interacted in an additive fashion. The unprecedented degree of synergy suggests that combining therapies directed at multiple targets can lead to breakthroughs in the treatment of complex metabolic disorders.

Although we obtained profound synergistic effects, a second-generation AAV vector, AAV2/5, was used in the current study. Newer generation AAV vectors (e.g., AAVrh10 or AAV2/9) mediate higher levels of transduction and have a broader distribution in the mouse brain (Foust et al., 2009; Hu et al., 2010). Therefore, they would be expected to increase efficacy even further. Another limitation in the current study is the fact that neonatal mice can only be exposed to sublethal, nonablative conditioning radiation (∼400 rads; Sands et al., 1993). Consequently, the triple-treated mice had donor hematopoietic chimerism of only 5–30%. It will be important to determine whether higher levels of engraftment significantly increases efficacy in this combination treatment paradigm. Finally, recent single-treatment studies using sophisticated approaches and the latest generation gene transfer vectors have reported median lifespans in the Twitcher mice of only 80–120 d (Gentner et al., 2010; Rafi et al., 2012). However, those approaches would likely increase treatment efficacy if used in combination.

Although this intensive therapeutic regimen is efficacious, it is not without side effects. In our experience, mice are particularly sensitive to 1-cycloserine in the weeks immediately following gene therapy and BMT. We observed a relatively high (30–50%) mortality before weaning if mice were injected with 50 mg/kg 1-cycloserine during this time. Nearly no mortality was observed if mice were given a dose of 25 mg/kg during the same period. This sensitivity was related to the combination of treatments, as it was not seen in animals treated with 1-cycloserine alone. Similarly, we have observed minimal mortality in animals treated with gene therapy, BMT or a combination of the two. It is likely that 1-cycloserine, which inhibits the synthesis of many glucosylated and galactosylated lipids in addition to psychosine, alters the normal balance of sphingolipids and reduces the ability of the brain to respond to the stress associated with conditioning radiation and intracranial injection of virus. The reduction of many sphingolipids and the potential for developmental damage highlights the need for a more specific inhibitor of psychosine synthesis, perhaps an inhibitor of galactosyltransferase. Although several other galactosylated neurolipids require the galactosyltransferase enzyme, fewer lipids would be disrupted compared with 1-cycloserine. Regardless, this does not diminish the magnitude of synergy observed when multiple defects are simultaneously targeted.

Even though HSCT carries a significant risk to the patient, it is the current standard of care for GLD. One limitation of HSCT is the slow turnover of hematopoietic-derived microglia in the brain. It has been shown previously that significant engraftment of donor-derived cells in the brains of mice was not observed until 160 d post-BMT (Kennedy and Abbkowitz, 1997), and there is nearly undetectable GALC activity in the brains of 36 d Twitcher mice treated with BMT at birth (Lin et al., 2007; Reddy et al., 2011; Hawkins-Salsbury et al., 2012b). Due to the aggressive
nature of GLD, it is likely that significant disease develops before a meaningful number of donor microglia can repopulate the CNS and provide therapeutic levels of GALC. Here we demonstrate that although triple-treated Twitcher mice have very few donor cells in the brain at 36 d of age (34 d post-transplant), the number of donor-derived (GFP-positive) cells increased in the brains of animals that were >160 d of age. This could be due to the fact that they lived long enough for significant numbers of donor cells to repopulate the brain. Alternatively, the long-lived animals may have had higher levels of hematopoietic reconstitution from an early age and thus a better long-term outcome. Regardless of the dynamics of CNS engraftment, once donor cells are in the brain, they can serve a dual role by contributing GALC to surrounding tissue and providing an immunomodulatory effect.

Although this therapeutic regimen dramatically increases lifespan and improves behavior, the mice eventually succumb to the disease. One likely explanation for the continued disease progression is that the distribution of the AAV vector, and thus GALC activity, is not sufficient to completely eliminate psychosine in various brain regions. In support of this hypothesis are the persistent, albeit low, levels of psychosine observed in the brains of triple-treated mice. Because psychosine is measured in whole-brain homogenates (or whole-nerve homogenates), it is possible that local concentrations of the lipid could be considerably higher in regions that are inaccessible to GALC. It is known that AAV vectors preferentially infect neurons as opposed to other non-neuronal cells, such as oligodendrocytes. In a disease such as GLD, which severely affects oligodendrocytes and white matter, this could be a significant limitation. Consistent with this hypothesis is the fact that, although myelin levels (LFB staining) appear relatively normal in 36-d-old treated Twitcher mice, the improved myelination does not persist, and myelin staining is dramatically decreased in 160-d-old treated Twitcher mice. Development of more potent or cell-specific vectors may significantly improve outcomes.

Another potential cause for the incomplete clinical response is the increasing neuroinflammation as the triple-treated mice age. Although the levels of various immune mediators in whole-brain homogenates were essentially normalized by triple therapy, CD68 and GFAP expression progressively increase. We also see a progressive increase in PAS-positive cells in the white matter tracts of treated Twitcher mice between 36 and 160 d of age. This apparent discrepancy could be due to localized increases in cytokine/chemokine expression that are masked by low levels of cytokines in relatively large areas of the brain that have little or no inflammation such as the cortex. Regional increases in cytokine/chemokine expression could be responsible for the localized increases in CD68 and GFAP staining and partial therapeutic response observed in triple-treated animals.
A prominent feature of GLD is the involvement of the peripheral nervous system. This pathology is reflected in the sciatic nerves of Twitcher mice, which show reduced myelin, decreased axonal density, increased edema, and infiltration of immune cells. Other than lowering psychosine, this triple-treatment regimen appears to have little effect on the disease in the sciatic nerve, which likely contributes to the continued clinical/behavioral decline. This is supported by the poor performance of the triple-treated mice in the wire-hang test, which measures limb strength and serves as a surrogate for the peripheral neuropathy. Interestingly, however, the sciatic nerves of long-lived triple-treated Twitcher mice have histological features referred to as “onion-bulb” formations. Onion-bulb formation has been observed previously in Twitcher mice following high-dose irradiation and BMT (Kagitani-Shimono et al., 2008). However, this histological feature appears to be more widespread in the triple-treated mice. Finally, the continued decline of the triple-treated Twitcher mice could be due to incomplete correction of the autonomic nervous system. It has been shown previously that the autonomic nervous system is affected in Twitcher mice and leads to thymic atrophy and peripheral lymphopenia (Galbiati et al., 2007).

Lysosomal enzymes, including GALK, are ubiquitously expressed. Therefore, deficiencies in these acid hydrolases would be expected to affect many, if not most cells of the body. Not surprisingly, this can activate multiple primary and secondary pathogenic mechanisms. In the case of GLD, targeting a single pathogenic mechanism results in modest to no efficacy. In contrast, synergistic increases in efficacy were observed when the primary defect was targeted with CNS-directed gene therapy and the secondary accumulation of psychosine and neuroinflammation was treated with l-cycloserine and BMT, respectively. It is likely that efficacy will be further increased if additional pathogenic mechanisms are targeted and/or when improved gene transfer vectors and more specific substrate inhibitors are developed. Although the specific parameters will need to be empirically determined on a case-by-case basis, this type of combination therapy provides a promising paradigm upon which to base treatments for this relatively large class of inherited metabolic diseases that are caused by both the primary genetic defect and multiple secondary consequences that affect multiple organ systems.

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