INTRODUCTION
N-Acetyl-glucosaminidase deficiency (Sanfilippo Syndrome type B, Mucopolysaccharidosis type IIIB) typically causes a pediatric onset disease characterized phenotypically, by progressive motor and cognitive deterioration, and histologically by accumulation of lyosomal inclusions in most tissues.¹ No current treatment is approved in humans. After the gene was identified,²,³ a murine model of mucopolysaccharidosis type IIIB (MPS IIIB) was created.⁴ The MPS IIIB mouse shares many of the biochemical, histological and clinical features with the human disease.⁴,⁵ Several groups have demonstrated the ability of either intracranial or systemic gene therapy approaches to reduce lysosomal distention in the brains of MPS IIIB mice.⁶⁻¹¹ Our group also demonstrated improvements in histology with corresponding improvements in neurologic function and life span using intracranial gene therapy.¹² Intracranial delivery has thus far demonstrated the most consistent improvement in disease progression. Increases of approximately 30% in life span have been observed with central nervous system (CNS)-directed therapies. An intracranial gene therapy approach is now being pursued in a larger animal model of MPS IIIB.¹³⁻¹⁵ Systemic-targeted gene therapy was shown to reduce lysosomal storage in peripheral organs. However, none of these single approaches completely eradicates intra-cytoplasmic inclusions or normalizes the disease phenotype. We previously attempted a combination of CNS-directed gene therapy and bone marrow transplantation with little or no benefit seen for the bone marrow transplantation arm. However, the level of chimerism was relatively low and toxicities from the radiation conditioning were evident in the transplant arm. In other lysosomal storage disease models, therapies to the CNS and the periphery, with enzyme replacement, bone marrow transplantation or gene therapy, have shown greatly improved disease correction especially when initiated in the neonatal period.¹⁶⁻¹⁹ Therefore, we hypothesized that neonatal combination therapy directed to both the CNS and the periphery would provide better correction of the disease, especially if higher systemic levels of N-acetyl glucosaminidase (NAGLU) activity could be attained. We describe here the benefits obtained from each mode of gene therapy and the synergistic effect of combining intracranial adeno-associated virus (AAV)-NAGLU and systemic lentiviral-NAGLU gene therapy.

RESULTS
Treatments
All gene therapy injections were performed in mice pups at 2–4 days of age. Intracranial AAV-NAGLU treatment was...
performed as described previously with six direct injections of 2 μl each into frontal, temporal and cerebellar regions of the brain of vector at a concentration of $1.5 \times 10^{12}$ viral particles per ml. Intravenous lentiviral-NAGLU injections were also performed as described previously by injection of 100 μl of $1.6 \times 10^6$ infectious units per ml viral aliquot into the superficial temporal vein.

NAGLU activity

Biochemical analysis of $N$-acetyl-glucosaminidase activity for various organs was determined in mice from each group at ~8 months of age and compared with untreated MPS IIIB animals (MPS IIIB NO TX; Figure 1). The MPS IIIB animals receiving only intravenous lentiviral vector treatment (MPS IIIB IV-LENTI) had detectable NAGLU activity in all organs assayed (<2% of normal in the brain ($P<0.05$) and kidneys (not significant (NS)), 11% in heart ($P<0.01$), 12% in lungs (NS), 34% in liver ($P<0.001$), 34% in spleen ($P<0.05$) and 28% in the serum (NS; data not shown). Conversely, animals treated only with intracranial AAV (MPS IIIB IC-AAV) had approximately 200% ($P<0.05$), 3% (NS) and 5% (NS) of normal activity in the brain, liver and serum (data not shown), respectively, and little or no activity in the spleen, heart, lungs or kidneys. Combination therapy (MPS IIIB BOTH) yielded NAGLU activity levels of 424% for brain ($P<0.01$), 13.6% for liver ($P<0.001$), 6.7% for heart ($P<0.01$), 19.8% for spleen ($P<0.01$), 2.9% for lungs ($P<0.05$), 42% for serum (NS) and <1% in the kidneys (NS). Normal mice treated with both therapies (NORMAL BOTH) had no significant change in NAGLU activity from normal mice for the visceral organs ($P>0.05$ for every comparison).

Secondary enzyme activity

In most lysosomal storage disorders, other lysosomal enzymes, such as $\beta$-glucuronidase (GUSB), are elevated. The resolution of

![Figure 1](image_url)

**Figure 1.** NAGLU activity was measured in brain, liver, heart, spleen, kidneys and lungs from untreated and treated MPS IIIB mice. Mean activity levels ± s.e.m. for each group relative to the Normal no treatment (NO TX) group are depicted in the graphs. BOTH, both IV-LENTI and IC-AAV treatments; IC-AAV, intracranial adeno-associated viral NAGLU vector; IV-LENTI, intravenous lentiviral NAGLU vector. *$P<0.05$; **$P<0.01$; ***$P<0.001$ compared with MPS IIIB NO TX group.
this secondary elevation has been used as a surrogate for therapeutic response. Secondary elevations of GUSB activity in the various groups are depicted in Figure 2 and statistically compared with the MPS IIIB NO TX group. Untreated MPS IIIB (MPS IIIB NO TX) animals had significant elevations in GUSB compared with normal mice for all organs assayed: brain (358%, $P < 0.001$), liver (175%, $P < 0.01$), spleen (343%, $P < 0.05$), heart (619%, $P < 0.05$), lungs (645%, $P < 0.001$) and kidneys (439%, $P < 0.001$). IC-AAV mice had significant reductions (nearly normal) in secondary elevations compared with untreated MPS IIIB mice in the brain ($P < 0.0001$), 387% normal in the lungs ($P < 0.01$), but no significant reductions in the other visceral organs. IV-LENTI-treated mice had significant reductions in GUSB activity in the brain (228%, $P < 0.01$), kidneys (217%, $P < 0.05$), and lungs (227%, $P < 0.0001$). Combination treatment resulted in normalization of levels in the brain (95%, $P < 0.0001$), and significant reductions in the kidneys (188%, $P < 0.01$) and lungs (183%, $P < 0.0001$). GUSB activity in the spleen and heart were reduced in the combination arm but did not reach significance ($P = 0.052$ and 0.099, respectively). The combination-treated wild-type animals remained within the normal range except for the brain, with a significant reduction to 82% of normal.

Histology

MPS IIIB BOTH mice had the best response in the CNS, with marked reduction in lysosomal storage in glial and meningeal cells as well as in cortical neurons as depicted in Figure 3. These same cell types in MPS IIIB IC-AAV mice had a less striking but definite response to the intracranial therapy. MPS IIIB IV-LENTI mice had no response in neurons and only mild reduction in storage in glial and meningeal cells (Table 1). The spleen and liver in treated mice...
showed minimal or no response with IC-AAV therapy only (Table 1). However, there was a clear and similar reduction in storage in these sites in both the IV-LENTI and BOTH groups. As a group, the mice treated with combined IC-AAV and IV-LENTI therapy had the best response with the greatest reduction in storage.

Glycosaminoglycan (GAG) analysis
In order to assess the lysosomal storage more quantitatively, the Sensi-Pro Non-Reducing End (NRE) assay was performed on brain homogenates of adult mice from each group in order to measure the pathologic GAG (pGAG) accumulation of heparan sulfate. Figure 4 depicts the reduction in brain heparan sulfate by each treatment. Compared with the MPS IIIB NO TX group, MPS IIIB IV-LENTI had no significant reduction in pGAG (85% of NO TX, \( P = \text{NS} \)), MPS IIIB IC-AAV (24% of NO TX, \( P < 0.01 \)) and MPS IIIB BOTH (18% of NO TX, \( P < 0.01 \)) had significantly reduced pGAG. Normal mice had no detectable pGAG.

Circadian activity
Long-term running wheel recordings from 14 to 24 weeks of age were performed with mice from each experimental group under a 12-h light:12-h dark cycle. We previously described a difference between untreated MPS IIIB and normal animals in two circadian measurements: percentage of activity occurring during the light phase (greater in MPS IIIB) and the time from lights off to activity

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**Table 1.** Lysosomal storage in brain and viscera by light microscopy in MPS IIIB mice after treatment with intracranial AAV, intravenous lentiviral or combined gene therapy

| Tissue          | IC-AAV (n = 4) | IV-Lenti (n = 4) | BOTH (n = 3) |
|-----------------|---------------|-----------------|-------------|
| Liver           | NC            | NC              | NC          |
| Hepatocytes     | NC            | NC              | NC          |
| Kupffer cells   | NC            | NC              | NC          |
| Spleen          | NC            | ↓↓↓↓↓↓↓↓↓↓      | ↓↓↓↓↓↓↓↓↓↓   |
| Sinus lining cells | NC↓↓↓↓↓↓↓↓↓↓     | ↓↓↓↓↓↓↓↓↓↓      | ↓↓↓↓↓↓↓↓↓↓   |
| Brain           | NC            | ↓↓↓↓↓↓↓↓↓↓      | ↓↓↓↓↓↓↓↓↓↓   |
| Cortical neurons | NC            | NC              | NC          |
| Meninges        | ↓↓↓↓↓↓↓↓↓↓     | ↓↓↓↓↓↓↓↓↓↓      | ↓↓↓↓↓↓↓↓↓↓   |
| Glia            | ↓↓↓↓↓↓↓↓↓↓     | ↓↓↓↓↓↓↓↓↓↓      | ↓↓↓↓↓↓↓↓↓↓   |

Abbreviations: IC-AAV, intracranial adeno-associated virus/S-N-acetyl glucosaminidase; IV-LENTI, intravenous lentiviral-N-acetyl glucosaminidase; MPS IIIB, mucopolysaccharidosis type IIIB; NC, no change in lysosomal storage from age-matched untreated MPS IIIB mouse. ↓↓↓↓↓↓↓↓↓↓, Slight reduction in storage; ↓↓↓↓↓↓↓↓↓↓, moderate reduction in storage; ↓↓↓↓↓↓↓↓↓↓, marked reduction in storage, all compared with age-matched untreated MPS IIIB mouse.

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*Figure 3.* Lysosomal inclusions in parietal cortex. Representative sections of parietal cortex of (a) MPS IIIB, (b) MPS IIIB IV-LENTI, (c) MPS IIIB IC-AAV (d) MPS IIIB BOTH mice are shown. Black arrows indicate neuronal lysosomal inclusions and white arrows indicate lysosomal distension in glial cells.

*Figure 4.* Heparan sulfate levels in the brain of untreated and treated mice. Pathologic glycosaminoglycan (pGAG) levels in brain homogenates + s.e.m. for each group are depicted in pg per mg of protein (\( N = 4–8 \) for each group, NS—not significant, **\( P < 0.01 \), ****\( P < 0.0001 \) compared with the MPS IIIB NO TX group).
onset (shorter in MPS IIIB). Similar differences were observed in the current study, however, percent activity during the light phase did not reach statistical significance (figure not shown) after correction for multiple comparisons between the MPS IIIB NO TX (mean = 15.14%) and the Normal NO TX group (mean = 9.60%; \( P = 0.14 \)). Although the average activity of the MPS IIIB IC-AAV (mean = 14.23%) and MPS IIIB BOTH (mean = 12.03%) groups were increasingly numerically improved from the MPS IIIB NO TX group and closer to the Normal NO TX group, they were not statistically significantly different after correction for multiple groups.

The time of daily activity onset (phase angle of entrainment) was significantly earlier in the MPS IIIB NO TX animals compared with Normal NO TX mice (\( P = 0.007 \); Figure 5). There was no difference in the phase angle of entrainment between the MPS IIIB NO TX and the MPS IIIB IV-LENTI groups. However, the MPS IIIB IC-AAV trended toward improvement (\( P = 0.078 \)) and MPS IIIB BOTH was significantly improved relative to the untreated MPS IIIB mice (\( P = 0.003 \)). The MPS IIIB BOTH group phase angle is not different than the Normal NO TX (\( P = 0.710 \)).

Untreated MPS IIIB and normal animals had no difference in total daily activity level similar to published results.5 No other treatment group was different than normal or MPS IIIB untreated mice for total activity.

The combination-treated normal animals were no different than the normal untreated animals for any circadian parameter tested.

Auditory function
Auditory-evoked brainstem response thresholds were performed on each group at approximately 8–8.5 months of age. Similar to our previous findings, MPS IIIB NO TX mice had significantly diminished hearing compared with normal animals, reaching the maximum sound output level of the equipment at several frequencies without generating a response (Figure 6). Each of the MPS IIIB treatment groups had a consistently lower response threshold at all frequencies tested compared with MPS IIIB NO TX mice. Based on a group-wise comparison of the treatment groups, there appeared to be a trend towards lower thresholds in the IC-AAV group compared with the IV-LENTI group. MPS IIIB BOTH mice were significantly improved compared with either IC-AAV or IV-LENTI mice but remains significantly different than the Normal NO TX group (\( P < 0.001 \)). Surprisingly, the Normal BOTH animals also had diminished hearing compared with Normal NO TX animals. This presumably reflects some adverse effect of virus administration on hearing, but the Normal BOTH animals still had demonstrably better hearing thresholds than MPS IIIB mice.

Motor function
We previously demonstrated IC-AAV treatment could delay the progression of motor deficits in MPS IIIB mice as measured using a rocking rotarod. In the current study, we demonstrate that each of the treatment interventions significantly delayed the progression of motor dysfunction compared with MPS IIIB NO TX animals (Figure 7). The median time to a latency of \( \leq 60 \) s on the rod increased from \( \sim 40 \) weeks in MPS IIIB NO TX animals to \( \sim 78 \) weeks in MPS IIIB BOTH mice. MPS IIIB IC-AAV mice reached the median latency at \( \sim 62 \) weeks and MPS IIIB IV-LENTI mice reached that latency at \( \sim 52 \) weeks. The time for the MPS IIIB BOTH mice to reach the median latency was significantly greater than any other MPS IIIB group. The median had not been reached for either Normal group at 90 weeks of age and is statistically longer than the MPS IIIB BOTH group (\( P = 0.05 \)).
Life span

We showed previously that IC-AAV treatment prolongs the life of MPS IIIB animals by >100 days. We find a similar increase in median life span of IC-AAV in this study (Figure 8). All treatment groups demonstrate a significant increase in median survival relative to MPS IIIB NO TX (322 days). MPS IIIB IV-LENTI median survival increased to 358 days, MPS IIIB IC-AAV to 452 days and MPS IIIB BOTH to 612 days. There are no apparent differences in survival between the Normal and Normal BOTH groups up to 720 days of age (data not shown), and median survival has not been reached in these groups, and is significantly longer than any MPS IIIB treatment group.

Adverse events

We had the opportunity to examine at least four mice from all groups at ~280 days of age and two MPS IIIB IC-AAV, two MPS IIIB BOTH and nine Normal BOTH mice from 588–726 days of age for tumor formation. None of the mice examined had demonstrable tumors on gross examination.

DISCUSSION

The most important measures of clinical benefit for any therapy are quality of life and quantity of life. We have chosen the neonatal timeframe for treatment because lysosomal storage disease treatment by bone marrow transplant in humans has demonstrated that the earliest possible treatment must be pursued to prevent irreversible neurologic damage. This is likely to be the most efficacious intervention timeframe in humans once newborn screening for this disorder becomes commonplace. We have demonstrated that neonatal treatment with either an intravenous injection of a lentiviral vector, intracranial injection of an AAV vector or the combination of both can significantly improve surrogates for quality of life (motor function, hearing and circadian rhythm) and increase life span. In all cases, the systemic approach was less efficacious than the intracranial approach. This is not surprising given the predominantly neurologic manifestations of MPS IIIB. However, the human disease has manifestations that are not in the CNS, such as hepatomegaly, hernias, diarrhea and ear infections with effects on hearing that may be a combination of central and peripheral neurologic effects and ear structure abnormalities.22–26 In addition, some of the benefit in the IV-LENTI group may be related to the very small amount of activity observed in the brains of these animals, presumable from a small amount of lentiviral transduction across the blood–brain barrier as has been seen by other investigators.27 This may be supported by the relatively small decrease in brain heparan sulfate in this group, which was not statistically significant. We also cannot rule out the possibility that greater benefit might be achievable from the systemic approach if we obtained higher expression of NAGLU by either using a higher or repeated dose of vector. Although we were able to attain supra-normal levels in the brain with the combination of systemic and intracranial treatments, the levels of NAGLU activity were less than half normal in all other organs assayed. Interestingly, in our study, the combination approach consistently yielded lower organ activity than the systemic monotherapy and higher brain activity than the CNS-directed monotherapy. This may be due to disruption of the blood–brain barrier during the CNS injections allowing systemic vector to shunt to the brain reducing the other organ viral exposure. We attempted to determine the relative contribution of lentiviral and AAV transduction in the brain. However, because of the presence of repetitive elements, identical inserts and poor DNA quality after homogenization neither the quantitative PCR nor in situ hybridization approaches to quantify AAV versus lentivirus were successful. We cannot rule out the possibility that the combination approach animals received a greater vector copy number than the IC-AAV group, despite the same injection volume from the same lot of virus, that could explain the differences between the groups.

We observed that some organs, such as lungs, heart and kidneys seem more resistant to transduction as has been observed by other gene therapy investigations,27 but still have substantial reduction of secondary enzyme elevation. This abrogation of secondary elevation may be through low-level NAGLU enzyme cross correction, from more highly transduced organs such as the liver or spleen. However, this low-level cross correction is not enough to markedly reduce lysosomal inclusions histologically. Perhaps the level of cross correction needed to reduce secondary enzyme elevations is lower than what is needed to markedly reduce lysosomal distention. This has been observed with the liver-directed gene therapy for MPS VII as well.17 The adverse effects on auditory-evoked brainstem response thresholds following virus administration in the Normal group most likely did not result from AAV as this group was not adversely affected in our prior study, which evaluated IC AAV given in the same manner. Whether the adverse effects are attributable to a direct effect of lentivirus or rather toxic effects of excess NAGLU to normal hearing is not yet known. However, this could be addressed by the determination of long-term effects on hearing structure and function after systemic lentiviral treatment of Normal mice with either NAGLU or a reporter enzyme.

As we originally hypothesized, the combination approach appears to have the greatest benefit. Although MPS IIIB is a simple monogenic disease, it has a complex biochemical, histological and clinical phenotype. As NAGLU is expressed in virtually every cell of the body, most tissues have some lysosomal storage. Consequently, multiple tissues must be treated to provide the maximum benefit. Interestingly, the combination therapy appears to be additive for some measures that affect quality of life such as hearing and motor function, and synergistic for measures of circadian function (difference in time to activity onset of 2 min with IV-LENTI, 22 min with IC-AAV and 38 min with BOTH) and overall survival (difference in median survival of 36 days with IV-LENTI, 130 days with IC-AAV and 290 days with BOTH). Not surprisingly, the combination treatment was not bested by either single therapy in any functional assessment. Unfortunately, despite long-term functional improvements and substantial benefits in survival of the combination approach, the progression of disease during the last several weeks of life for each of the
groups remained similar with poor coat care, urinary retention and loss of balance.

The combination therapy approach using two different vectors in the current study is not the only dual approach that can be envisioned. We selected a lentiviral vector for the systemic approach for its efficacy at producing sustained expression of a protein product and the lack of known immune inactivation in human trials. A higher sustained level of NAGLU production may have been obtained with lentiviral transduced bone marrow selected for high enzyme level as has been demonstrated by others. Alternatively, a systemic AAV9 vector approach may yield similar systemic benefits with a single peripheral injection site; however, this serotype has not yet been used in humans. In contrast to lentivirus trials, human systemic AAV gene therapy approaches have been limited by immune responses to the virus-infected cells. The systemic AAV approach needs further study to avoid immune inactivation that has been seen in human trials.

In addition to determining the efficacy of a dual gene therapy approach, the safety of AAV and lentiviral vectors requires additional study. Several recent studies have demonstrated the long-term safety of systemic delivery of AAV vectors in both rodents and non-human primates. However, a number of studies have also shown an increase in tumorigenesis in AAV-treated animals.

In summary, combination neonatal intracranial and systemic NAGLU gene therapy provides significant and clinically meaningful therapeutic benefit in a mouse model of MPS IIIB. However, study of additional interventions is warranted as the current strategy is not completely corrective.

**MATERIALS AND METHODS**

**Viral constructs**

AAV-NAGLU was constructed as previously described with a huNAGLU cDNA (gift of Elizabeth Neufeld), AAV2 genome, CMV enhancer, chicken β-actin promoter, SV40 poly-A signal and 3′ untranslated region from the rabbit 1-globin gene. Vector was produced at the University of Florida Vector Core with a pseudotype AAVs capsid, and diluted to 1.5 x 10^11 virus particles per ml with lactated Ringer’s solution before freezing at 85°C.

Lentiviral-NAGLU was constructed with the same huNAGLU cDNA into the MND vector plasmid (kind gift from D Kohn) with a delta-gag, central polyurine tract and the myeloproliferative sarcoma virus enhancer, negative control region deleted, D587rev primer-binding site-substituted (MND) promoter, SV40 poly-A tail and delta u3 3′long terminal repeat. Vector was produced in the Sands’ lab with a four-plasmid system in 293T cells and concentrated to 1.6 x 10^7 infectious units per ml before aliquoting and freezing at 85°C.

**Mice**

C57BL/6 NAGLU-deficient mice (kind gift from E Neufeld) were maintained by strict sibling mating by MSS at Washington University School of Medicine. Genotyping was done on tissue of newborn mice by enzyme assay or NAGLU exon 6 and neomycin insertion cassette PCR. All procedures on animals were in accordance with the Guidelines of Institutional Animal Care and Use Committee at Washington University in St Louis.

**Treatments**

At 2–4 days of age all mice were allocated to treatment groups: untreated MPS IIIB (MPS IIIB NO TX, n = 19, 15 males), MPS IIIB treated with intracranial AAV-NAGLU (MPS IIIB IC-AAV, n = 19, 10 males), MPS IIIB treated with intravenous lentiviral NAGLU (MPS IIIB IV-LENTI, n = 19, 7 males), MPS IIIB treated with both intracranial AAV-NAGLU and intravenous lentiviral NAGLU (MPS IIIB BOTH, n = 16, 9 males), male untreated (Normal NO TX, n = 15, 10 males) and normal with both intracranial AAV-NAGLU and intravenous lentiviral NAGLU (Normal BOTH, n = 17, 8 males). All gene therapy injections were performed in mice pups at 2–4 days of age. Intracranial AAV-NAGLU treatment was performed as described previously with six direct injections of 2 µl each into frontal, temporal and cerebellar regions of the brain using a 32-G needle. Intravenous lentiviral-NAGLU injections were also performed as described previously by injection of 100 µl of viral aliquot into the superficial temporal vein. When combined, the AAV injections were performed first and the systemic injections were performed within 5–60 min.

**Histology and biochemistry**

Mice from each group (n = 3–8 for MPS IIIB NO TX and MPS IIIB BOTH, and n = 4–8 all other groups) between 242 and 259 days of age were killed by CO2 asphyxiation. Liver, spleen, kidneys, heart, lungs and brain were harvested. Part of each organ was immersion fixed in 2% glutaraldehyde/4% paraformaldehyde in phosphate-buffered saline and part was flash frozen in liquid nitrogen and stored at −85°C until mechanical homogenization in 10 mTris (pH 7.5), 150 mNaCl, 1 mDithiothreitol and 0.2% Triton X-100. Fixed tissue was embedded in Spurr’s resin and 1-µm-thick sections were stained with toluidine blue before blinded evaluation of lysosomal storage and vacuolization.

Cell debris was pelleted and supernatants were collected for enzyme assays of NAGLU and GUSB activity. Duplicate NAGLU assays were performed using 20 µl of supernatant added to 40 µl of 0.2 m4-methylumbellifere-N-acetyl-β-D-glucopyranosidase (Sigma, St Louis, MO, USA), 0.1 mNaC2H3O2, 0.5 mM ml-1 bovine serum albumin and incubated at 37°C. Reactions were stopped with 1 ml of 0.2 mNa2CO3, 0.32 m glycine. Substrate cleavage was determined at excitation 365 nm and emission 448 nm using a Hitachi F-2000 Flourescence Spectrophotometer using a standard curve of 0.5–5 nm ml-1. Specific activity was corrected for protein concentration. Duplicate GUSB assays were performed similarly using the 4-methylumbellifere-N-acetyl-β-D-glucopyranosidase enzyme assay method previously described.

Activity levels were analyzed by one-way analysis of variance (ANOVA) for comparison of treatments to NO TX groups after confirmation of a difference between NORMAL and MPS IIIB NO TX groups by Student’s t-test. Brown-Forsythe Test of equal variances was satisfied as not significant for organ comparisons of brain, liver, heart, lungs, spleen and kidneys of GUSB.

**GAG analysis**

Homogenized brain samples (N = 4–8 for each group, all from mice older than 200 days) were coded with a numerical identifier and sent blinded to Zacharon Pharmaceuticals Inc. (San Diego, CA, USA) for pGAG analysis using Sensi-Pro NRE assay. pGAG are the GAG fragments present due to the deficiency of the specific lysosomal enzyme. The Sensi-Pro NRE assay is a highly specific and sensitive assay that uses high-performance liquid chromatography to quantitate the reduction in lysosomal GAG accumulation, by labeling and quantifying the NREs of these GAG fragments. In MPSIIIB, the pGAG markers are unique NRE-derived trisaccharides that enhance the efficacy of both neuronal stem cell- and hematopoietic stem cell-mediated transplantation in the murine model of Sandhoff disease. The caveat to this approach is the relatively high morbidity and mortality of stem cell transplant methods in humans from the conditioning regimens, infections, and graft-versus-host transplants from graft-versus-host disease.

In conclusion, combination neonatal intracranial and systemic NAGLU gene therapy provides significant and clinically meaningful therapeutic benefit in a mouse model of MPS IIIB. However, study of additional interventions is warranted as the current strategy is not completely corrective.
running wheel within light-tight ventilated chambers illuminated by fluorescent bulbs (F30T12-SP41-RS, General Electric, Fairfield, CT, USA, 3.9 × 10^17 to 6.9 × 10^18 photons s⁻¹ m⁻²) at the bottom of the cages. Wheel running activity was recorded in 1 min bins (Clocklab, Actimetrics, Evanston, IL, USA), whereas mice were exposed to a light-dark schedule (lights on at 07:00 and off at 19:00 hours).

We analyzed phase angle of entrainment (delay between daily light offset and onset of activity), total daily activity and proportion of daily activity in the light phase of the photocycle using Clocklab. Statistical analysis was by repeated-measures ANOVA.

Auditory evaluation

Auditory-evoked brainstem responses were performed at 8–8.5 months of age using procedures similar to those described previously. Mice were anesthetized with ketamine/xylazine (85/15 mg kg⁻¹), i.p. while core temperature was maintained at 37.0 ± 1.0 °C by a thermostatically regulated heating pad monitored via a rectal probe (Yellow Springs). Platinum needle electrodes (Grass, West Warwick, RI, USA) were placed subcutaneously in the back, vertex and behind the right ear while connecting to a Grass P15 differential amplifier (100–10,000 Hz, 100 ×). A Cambridge Electronic Design Micro 1401 (Cambridge Electronic Design, Cambridge, UK) running SIGNAL and custom averaging software digitized the signal at 30 kHz. Toneburst stimuli at 5, 10, 20, 28.3 and 40 kHz were delivered 1000 times at 20 s⁻¹ using an Alpine SPS-ORCA coaxial speaker (Crutchfield, Charlottesville, VA, USA) located 10 cm lateral to the right ear. At each test frequency, the stimulus level was reduced in 5 dB minimum steps to determine the minimum sound pressure level required for visual detection of Wave I. Repeated-measure ANOVA with Dunnett’s multiple comparison correction was performed to compare each treatment group to the MPS IIIB NO TX control. Separate one-way ANOVA was performed for data from all groups at each test frequency, followed by Bonferroni multiple comparison tests. Sample sizes by group were as follows: MPS IIIB NO TX, n = 10; MPS IIIB IC-AAV, n = 9; MPS IIIB IV-LENTI, n = 9; MPS IIIB BOTH, n = 10; Normal NO TX, n = 10; Normal BOTH, n = 10.

Motor function assessment

As we described previously in this model, mice were trained on a rotarod (UGO Basile, Varese, Italy) moving at a speed of 10 r.p.m. that switched direction after each full rotation for up to 180 s per attempt, with three attempts per day. Tests were done every 28 days from 196 up to 672 days of age (MPS IIIB NO TX, n = 6; MPS IIIB IC-AAV, n = 9; MPS IIIB IV-LENTI, n = 10; MPS IIIB BOTH, n = 10; Normal NO TX, n = 9; Normal BOTH, n = 8). Longest latency to fall from the rotarod of the three attempts was used for comparisons. Data were interpreted using a Log Rank test for time to latency of <60 s on rotarod.

Life span

All treated animals were analyzed by intention to treat with a Gehan–Breslow–Yates test. Kaplan–Meier curves were generated to assess the effect of treatments on survival.

CONFLICT OF INTEREST

Jillian R Brown and Brett E Crawford are full-time employees and shareholders of Zacharon Pharmaceuticals. All other authors have no conflict of interest to report.

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