Systemic Delivery of AAVB1-GAA Clears Glycogen and Prolongs Survival in a Mouse Model of Pompe Disease

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Pompe disease is an autosomal recessive glycogen storage disorder caused by deficiency of the lysosomal enzyme acid alpha-glucosidase (GAA). GAA deficiency results in systemic lysosomal glycogen accumulation and cellular disruption in muscle and the central nervous system (CNS). Adeno-associated virus (AAV) gene therapy is ideal for Pompe disease, since a single systemic injection may correct both muscle and CNS pathologies. Using the Pompe mouse (B6;129-GaaTm1Rabn/J), this study sought to explore if AAVB1, a newly engineered vector with a high affinity for muscle and CNS, reduces systemic weakness and improves survival in adult mice. Three-month-old Gaa–/– animals were injected with either AAVB1 or AAV9 vectors expressing GAA and tissues were harvested 6 months later. Both AAV vectors prolonged survival. AAVB1-treated animals had a robust weight gain compared to the AAV9-treated group. Vector genome levels, GAA enzyme activity, and histological analysis indicated that both vectors transduced the heart efficiently, leading to glycogen clearance, and transduced the diaphragm and CNS at comparable levels. AAVB1-treated mice had higher GAA activity and greater glycogen clearance in the tongue. Finally, AAVB1-treated animals showed improved respiratory function comparable to wild-type animals. In conclusion, AAVB1-GAA offers a promising therapeutic option for the treatment of muscle and CNS in Pompe disease.

Keywords: Pompe disease, AAVB1, AAV9, respiratory, glycogen

INTRODUCTION

Pompe disease is an autosomal recessive lysosomal storage disorder caused by mutations in the GAA gene encoding acid alpha-glucosidase (GAA)—a lysosomal enzyme responsible for the hydrolysis of glycogen to glucose. Deficiency in GAA results in accumulation of glycogen in lysosomes and subsequent cellular dysfunction in cardiac, skeletal, and smooth muscles as well as in the central nervous system (CNS).1-6 Pompe disease can present early in life as infantile onset Pompe disease (IOPD) or later in childhood to adulthood as late onset Pompe disease (LOPD). IOPD patients have little to no GAA activity and, if left untreated, typically develop cardiorespiratory failure within the first year of life.7 LOPD patients have some residual GAA activity resulting in a more variable presentation of muscle weakness and respiratory insufficiency.8-11 Respiratory failure is a prominent cause of death in both types of Pompe disease.2,11 The only Food and Drug Administration (FDA)-approved treatment for Pompe disease is enzyme replacement therapy (ERT). However, systemically administered GAA does not cross the blood–brain barrier and therefore cannot treat the CNS pathology and affected respiratory motor neurons. Furthermore, ERT only partially corrects skeletal muscle abnormalities as a result of low uptake into muscle fibers. Consequently, two-thirds of IOPD patients eventually require ventilatory support,12

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and respiratory insufficiency persists among LOPD patients.\textsuperscript{13,14}

Gene therapy using adeno-associated virus (AAV) vectors is ideal for Pompe disease, since it is a monogenetic disorder. Contingent on the serotype and promoter, AAV gene therapy has the potential to treat both the muscle and motor neuron pathologies following a single administration. AAV gene therapy for Pompe disease has been extensively studied in the Gaa\textsuperscript{−/−} mouse model.\textsuperscript{15–25} These studies resulted in a Phase I/II clinical trial using AAV1-GAA intramuscular injections into the diaphragm of Pompe patients.\textsuperscript{26} This trial was the first to show that AAV1-GAA delivered to the diaphragm was safe and resulted in therapeutic benefits. Patients had increased unassisted tidal volumes and increased tolerance of unassisted breathing, with continued respiratory benefit seen 1 year post injection (p.i.).\textsuperscript{27} Furthermore, a clinical trial using AAV9-GAA injected intramuscularly into the tibialis anterior is currently underway to test the feasibility of gene therapy re-dosing.\textsuperscript{28}

Systemic delivery of AAV9-GAA in Gaa\textsuperscript{−/−} adult mice has yielded variable results, with some improvements in respiratory outcome measures but with variable glycogen clearance in muscle.\textsuperscript{17} An ever-evolving area within the gene therapy field is enhancing vector biology for more efficient transduction of disease relevant tissues. Many new capsid variants have emerged over the years by rational design or library screening. Thus, this study sought to explore novel vectors in order to optimize muscle and motor neuron transduction for Pompe disease. One of these new capsids, AAVB1, generated by library selection showed enhanced muscular and neuronal transduction compared to AAV9.\textsuperscript{29} In addition, this vector is more favorable for clinical application due to a reduced immune profile.\textsuperscript{26} Since muscle and CNS transduction is more efficient with AAVB1, it was hypothesized that AAVB1-GAA would be a more efficient therapy for overall glycogen clearance in Pompe disease.

**METHODS**

**Animals**

All experimental procedures were approved by the Institutional Animal Care and Use Committee at the University of Massachusetts Medical School. B6;129-Gaa\textsuperscript{TmIRabn/J} mice (stock number 004154) were obtained from Jackson Laboratory (Bar Harbour, ME) and were fed a standard mouse chow ad libitum. The mice were bred in-house by breeding B6;129-Gaa\textsuperscript{TmIRabn/J} males to B6;129-Gaa\textsuperscript{TmIRabn/J} females; animals heterozygous at the Gaa allele were used to create littermate controls. Prior to enrollment in the study, littermates were genotyped using standard polymerase chain reaction (PCR) methods to confirm the lack of Gaa gene.\textsuperscript{30,31} Adult male Gaa\textsuperscript{−/−} (n = 36) and wild-type (WT) controls (n = 10) were enrolled in the study at 3 months of age. Gaa\textsuperscript{−/−} mice were randomly and blindly assigned to one of three treatment groups: AAV9 (n = 10), AAVB1 (n = 10), or phosphate buffered saline (PBS; n = 10). Six additional animals were enrolled in the PBS group because of the high death rate in this group and the increased number of animals needed for the physiological studies. All WT animals were injected with 200 $\mu$L of PBS.

**AAV vectors**

AAV9 and AAVB1 vectors were generated using previously published methods.\textsuperscript{29} Vectors were purified by iodixanol gradient centrifugation followed by buffer exchange, as described previously.\textsuperscript{28} Final formulations of AAVB1-DES-hGAA and AAV9-DES-hGAA were in PBS. All vectors were generated and titered at the University of Massachusetts in the laboratory of Dr. Sena-Esteves using previously published methods.\textsuperscript{29} Either AAV9 or AAVB1 (1 × 10\textsuperscript{12} vector genomes [vg]) were injected in a total volume of 200 $\mu$L (or PBS for controls) into the tail vein of the mice.

**Behavior testing**

Four-limb grip strength was determined by a blinded investigator by measuring peak force using an Almenō\textsuperscript{®} digital grip strength meter (Ahlborn, Holzkirchen, Germany) equipped with a mesh screen. Mice were placed on the metal grid until they gripped it, and then they were gently pulled by the tail until they could no longer hold the grid. The grip force test was repeated twice, with a 15 min rest period in between, and the maximum force was recorded.

Inverted screen testing was performed by an investigator blinded to the treatment group or genotype on an apparatus, as described previously.\textsuperscript{32} Animals were placed on a square wire mesh of 30 cm\textsuperscript{2} with 25 mm\textsuperscript{2} holes, over a cushioned surface, which was rotated 180°. Animals were assessed for latency to falling for up to 2 min. This was repeated twice per session, with a 15 min rest period in between, and the longest hang time was recorded.

**Ventilation**

These studies determined the impact of systemic vector injection on ventilation. Ventilation was quantified 180 days p.i. using whole-body plethysmography in unrestrained, unanesthetized mice,
as previously described. An investigator blinded to the treatment group or genotype performed these studies. Awake, unrestrained mice were placed inside a 3.5” × 5.75” Plexiglas chamber (SCI REQ, Montreal, Canada), and data were collected in 10 s intervals. The Drorbaugh and Fenn equation was used to calculate respiratory volumes, including tidal volume and minute ventilation. During both a 30 min acclimation period and subsequent 60–90 min eupnea period, mice were exposed to normoxic air (FiO₂: 0.21; nitrogen balance). Then, the mice underwent a 10 min hypercapnic challenge (FiO₂: 0.21; FiCO₂: 0.07; nitrogen balance).

**Measures of airway resistance**

Pulmonary mechanics were performed at study endpoint using forced oscillometry (FlexiVent system, SCIREQ) at baseline and in response to incremental doses of methacholine, as previously described. In brief, animals were anesthetized with an intraperitoneal injection (i.p.) injection of a mixture of ketamine (90 mg/kg; Animal Health International) and xylazine (4.5 mg/kg; Propharma), and a tracheotomy was performed followed by insertion of pre-calibrated cannula into the trachea. Spontaneous respiratory effort was prevented using a neuromuscular blocking agent (pancuronium bromide: 2.5 mg/kg; Hospira, Inc., Lake Forest, IL). Respiratory mechanics were obtained and calculated using FlexiWare software (SCI REQ), as previously described. After the initial mechanical scan protocol, incremental doses (3.125, 6.25, 12.5, and 25 mg/mL) of methacholine were nebulized, as previously described. Between each dose, it was confirmed that the mouse was still alive by assessing the heartbeat.

**Vector genomes**

gDNA was extracted using a DNeasy blood and tissue kit (Qiagen, Valencia, CA) from the tissues harvested from AAV- and PBS-injected animals 180 days after injection. gDNA concentrations were determined using a Nanodrop 2000c (Thermo Fisher Scientific, Carlsbad CA). Recombinant AAV genome copies in the gDNA were quantified by quantitative PCR with an Applied Biosystems Vii7 (Life Technologies, Carlsbad, CA), as previously described. Briefly, DNA samples were assayed in triplicate, using primers and probe designed to amplify a sequence from the SV40 poly-A signal in the DES-hGAA vector (forward: AGCAATAGCATCACAATTTCAC; reverse: GCAGACATGTAAGATACATT-GATGAGTT; and probe: 6FAM-AGCATTTTTTTCACTGATTCTAGTTGTGGTTT GTC). A standard curve was generated using plasmid DNA containing the same SV40 poly-A target sequence.

**Histology**

The following tissues were harvested at 180 days after injection immediately following the final respiratory mechanics test: the medulla and spinal cord, heart, tongue, diaphragm, gastrocnemius muscle, and trachea.

GAA staining. The heart, gastrocnemius, tongue, diaphragm, medulla, and spinal cord tissues were extracted and post fixed by immersion in 4% paraformaldehyde (n = 4 animals per group) for 24 h and then transferred to 30% sucrose for cryoprotection until sinking. The brain stem and spinal cord were then blocked into the brain stem, cervical spinal cord, thoracic spinal cord, and lumbar spinal cord. Samples were embedded in optimal cutting temperature (OCT) compound and frozen. Sections (8 μm) were cut for the heart, gastrocnemius, tongue, and diaphragm; 40 μm sections were cut for the medulla and spinal cord. Sections stained by immunohistochemistry (IHC) for GAA, as previously described. Briefly, the tissue was incubated overnight in primary antibody against hGAA, 1:2,000 (rabbit polyclonal anti-hGAA antibody; Covance, Emeryville, CA). The following day, the tissue was washed in PBS, incubated in a biotinylated anti-rabbit IgG secondary antibody, 1:2,000 (rabbit polyclonal anti-hGAA antibody; Covance, Emeryville, CA). The following day, the tissue was washed in PBS, incubated in a biotinylated anti-rabbit IgG secondary antibody, 1:2,000 (rabbit polyclonal anti-hGAA antibody; Covance, Emeryville, CA) and coupled with a Vectastain ABC Kit and DAB for bright field microscopy.

Periodic acid–Schiff staining. The heart, tongue, diaphragm, trachea, and gastrocnemius muscle were either fixed in glutaraldehyde and then embedded in epoxy resin or immediately frozen in OCT. Tissues were processed, as previously described, for epoxy resin embedding. Polymerized blocks were trimmed and cut at 1.5 μm on a Leica Ultracut E ultramicrotome and mounted on charged slides. To evaluate the histological glycogen load, periodic acid–Schiff (PAS) staining was used, as previously described. Briefly, tissues were incubated in 1% periodic acid for 10 min at 60°C, rinsed with tap water for 1 min, and then immediately stained with Schiff’s reagent (Sigma–Aldrich, St. Louis, MO) at room temperature for 10 min, followed by a tap-water rinse until the water ran clear. Once dried completely, slides were coverslipped with Permount mounting medium (Thermo Fisher Scientific, Hampton NH) without dehydration in ethanol and the use of xylene. For the fresh-frozen tissue, 8 μm sections were stained with PAS according to standard
motor neurons, but because it has high activity in cardiac, skeletal, and smooth muscle as well as in motor neurons, but limited activity in the liver.\textsuperscript{37} Gaa\textsuperscript{−/−} mice and WT mice were injected with PBS as controls. Several parameters were assayed at 30, 90, and 180 days p.i. The experimental endpoint was at 180 days when the animals were approximately 9 months old. By 180 days p.i., 50\% of PBS-treated Gaa\textsuperscript{−/−} controls had survived, with the rest of the animals spontaneously dying between 76 and 173 days (Fig. 1A). Survival significantly improved in both the AAVB1- and AAV9-treated animals (p = 0.0037; Fig. 1A). However, only the AAVB1-treated animals displayed robust weight gain over time in comparison to the AAV9- and PBS-treated Gaa\textsuperscript{−/−} mice (Fig. 1B). WT animals were significantly heavier than all three Gaa\textsuperscript{−/−} groups at 30 days (p < 0.02). However, by 90 and 180 days, the difference in weight was only significant when comparing the WT group to both AAV9- and PBS-treated groups (p < 0.04). At 90 days, the AAVB1-treated Gaa\textsuperscript{−/−} animal weights were not significantly different from WT (p = 0.16), and by 180 days, their weights had significantly increased (p < 0.05), reaching near WT weights (p = 0.92).

Next, the study looked at the muscle strengths of treated and control groups. At 180 days p.i., AAV9- and PBS-treated groups had significantly lower four-limb strength (Fig. 1C) and inverted screen hang time (Fig. 1D) compared to WT, whereas AAVB1-treated animals were comparable to WT animals (p = 0.24) in four-limb strength (Fig. 1C). At all time points, Gaa\textsuperscript{−/−}-untreated and AAV9-treated mice had significantly lower inverted screen hang times (Fig. 1D) than WT controls, whereas AAVB1-treated animals were able to maintain strength at near WT levels at 30 and 90 days p.i.

**Vector genome copies**

In order to compare the transduction of AAVB1 to AAV9, vector genome copies were measured in respiratory-specific tissues, peripheral tissues, and the spinal cord. In the respiratory tissues (Fig. 2A), a trend of increased transduction of AAVB1 compared to AAV9 was observed in the diaphragm and lung, neither of which was statistically significant. Significantly greater levels were, however, measured in the tongue base, which includes the genioglossus muscle—a muscle necessary for maintaining airway patency. In contrast, there was lower transduction of AAVB1 compared to AAV9 in the trachea (p = 0.3 trachea). In the medulla, cervical, and lumbar spinal cord (Fig. 2B), there was similar transduction between the two groups, but a significantly greater transduction was observed with AAVB1 in the thoracic spinal cord (p = 0.001). Genome copy numbers were identical
for both AAV vectors in the heart, gastrocnemius muscle, and liver (Fig. 2C).

**GAA immunohistochemistry**

IHC for hGAA protein was performed in order to assess vector expression in AAVB1- and AAV9-treated animals 180 days p.i. Cross-sections of cardiac, respiratory, and hind-limb muscles (Fig. 3A), as well as sections of the spinal cord (Fig. 3B), were stained with an anti-hGAA antibody, specific for human GAA produced from the vector. Robust GAA expression was evident in the cardiac muscle of both the AAVB1- and AAV9-treated animals. Expression in the diaphragm and gastrocnemius muscle occurred, but this was weaker than in the heart. Superior expression in the tongue was observed in AAVB1-treated animals compared to AAV9. In sections of the spinal cord from the medulla, cervical, and thoracic regions, motor neurons were transduced in both AAVB1- and AAV9-treated animals.

**GAA activity assay**

To assess the amount of functional GAA enzyme present, GAA enzyme activity assays were performed 180 days p.i. Supraphysiological levels of GAA enzyme activity were observed in the heart, diaphragm, tongue, gastrocnemius muscle, and lung but not in liver from both AAV-treated groups (Fig. 4A–F). GAA activity was comparable between AAVB1- and AAV9-treated animals in all muscles, but only in the heart was it significantly elevated compared to that in WT animals (\( p < 0.05 \) for both groups; Fig. 4A). Lung GAA activity in AAVB1-treated animals was significantly higher than in WT controls (\( p = 0.03 \); Fig. 4E). In contrast, the liver GAA levels in AAV-treated animals were considerably below WT, most likely due to the use of the tissue-restrictive desmin promoter (Fig. 4F). GAA activity was below the detection limit of the enzyme assay in the trachea, medulla, cervical, thoracic, and lumbar spinal cord of both AAV treatment groups (data not shown).

**Glycogen clearance**

Next, the study examined the impact of AAV treatment on lysosomal glycogen accumulation by histological analysis of glycogen abundance in sections of respiratory, cardiac, and hind-limb muscles stained with PAS. Reduced glycogen accumulation...
was observed in each of the muscles studied from both AAVB1- and AAV9-treated mice. A nearly equal, robust clearance of glycogen storage was observed in both the heart (Fig. 5A) and diaphragm (Fig. 5B) in animals injected with either AAV. However, in AAVB1-treated animals, glycogen clearance was greater in the tongue (Fig. 5C), whereas clearance in the gastrocnemius appeared greater in AAV9-treated animals (Fig. 5D). Quantification of clearance of glycogen was determined by biochemical glycogen assay (Fig. 5E–H). Near-complete clearance of glycogen was observed in the heart, mirroring the histology (Fig. 5A), with significant differences when comparing PBS-treated Gaa−/− mice groups to WT, AAV9, and AAVB1, and both treatment groups were not statistically significant from WT (Fig. 5E). Similarly, within the gastrocnemius and diaphragm, significant reduction of glycogen was observed in all groups when compared to PBS-treated Gaa−/− mice (Fig. 5F and 5H). In the tongue, glycogen clearance was pronounced in AAVB1-treated animals, similar to the observation of PAS staining, and glycogen accumulation was significantly decreased from PBS-treated Gaa−/− mice and not significantly different from WT (Fig. 5G). In contrast, in the tongues of AAV9-treated animals, glycogen accumulation was significantly increased over WT and not significantly different from PBS-treated Gaa−/− mice.

Respiratory outcome measures

The impact of gene therapy on breathing and lung mechanics was examined over time. Using whole-body plethysmography, lung volumes and ventilation parameters were tested during eupnea (normoxic conditions: FiO2: 0.21; nitrogen balance) and during a respiratory challenge (hypercapnic conditions: FiO2: 0.21; FiCO2: 0.07; nitrogen balance). No significant differences occurred between groups at baseline, 30 days p.i., or 90 days p.i. (data not shown). Thus, the impact on breathing was evaluated at 180 days p.i. as a percent change during respiratory challenge over baseline breathing (Fig. 6A–G). Statistically significant differences between WT and Gaa−/− mice were observed at 180 days under hypercapnic conditions, specifically in tidal volume (TV; Fig. 6B; \( p = 0.02 \)), minute ventilation (MV; Fig. 6C; \( p < 0.05 \)), peak inspiratory flow (PIF; Fig. 6D; \( p = 0.001 \)), and inspiratory time (Ti; Fig. 6F; \( p = 0.01 \)). PIF is a reflection of diaphragm and inspiratory muscle strength, and a higher Ti is a reflection of increased upper airway resistance. In addition, AAV9-treated animals showed reduced TV (Fig. 6B) and PIF (Fig. 6D) in response to a respiratory challenge compared to WT animals (\( p < 0.05 \) and \( p = 0.01 \), respectively), but AAVB1 animals did not. No significant differences in MV or Ti were observed between AAV9 and WT (\( p > 0.09 \)) or AAVB1 and WT animals (\( p > 0.44 \)). In addition, AAVB1-treated animals had a more robust decline in Ti during a respiratory challenge.
Figure 3. Robust acid alpha-glucosidase (GAA) expression was noted by immunohistochemistry in both AAV9 and AAVB1 groups. Tissues were harvested 180 days following systemic AAV9 and AAVB1 injections, and the presence of GAA was detected by immunohistochemistry (brown). (A) GAA expression was seen in the heart, diaphragm, tongue, and gastrocnemius muscle. (B) In the medulla and spinal cord, motor neurons in both AAV9- and AAVB1-injected animals stained positive for GAA. Scale bar = 100 µm. Color images available online at www.liebertpub.com/hum

Figure 4. GAA enzymatic activity evident 6 months following AAV injection. (A) GAA activity in both AAV9- and AAVB1-injected Gaa–/– mice was robust in the heart, with levels well above WT levels. (B–D) GAA activity was not significantly different between AAV9 and AAVB1 animals in the diaphragm (B), tongue (C), and gastrocnemius muscle (D). (E) Lung GAA activity was significantly higher in the AAVB1-treated group compared to the AAV9 group. (F) Liver GAA activity levels were significantly lower in both AAV groups compared to WT. Data are represented as mean ± SEM. N.D., not detected. *p < 0.05.
challenge than PBS-treated Gaa<sup>−/−</sup> animals (Fig. 6E; p = 0.04).

As has been shown, robust accumulation of glycogen in the airway smooth muscle occurs in the Gaa<sup>−/−</sup> mouse model. This results in disruption of pulmonary mechanics, namely an inability of the airways to react to a bronchoconstrictive agent (i.e., methacholine).<sup>36</sup> Consistent with these findings, when a methacholine challenge was performed on sedated, intubated, and ventilated WT and Gaa<sup>−/−</sup> mice, significant differences in tissue and small airway resistance was seen between WT and Gaa<sup>−/−</sup> mice at a methacholine dose of 12.5 mg/mL and 25 mg/mL (Fig. 6H). However, no difference was observed in resistance between PBS-treated Gaa<sup>−/−</sup> animals and either AAV9- or AAVB1-treated groups (p = 0.55 and p = 0.82, respectively). Moreover, both AAV- and PBS-treated Gaa<sup>−/−</sup>
animals had significant airway hyporesponsiveness to methacholine when compared to WT animals (Fig. 6H). This finding was consistent with histological evidence showing continued glycogen accumulation in trachea of AAV9- and AAVB1-treated animals (Fig. 6I and J, respectively).

**DISCUSSION**

The most important finding in this study is that a single systemic injection of either AAVB1-DES-hGAA or AAV9-DES-hGAA significantly prolongs survival of Pompe (B6;129-Gaa<sup>Tm1Rabn</sup>/J Gaa<sup>−/−</sup>) mice. Both vectors robustly transduced the heart, increased enzyme activity, and cleared glycogen in the myocardium. This cardiac transduction is the most likely explanation for the increase in survival. Furthermore, based on these studies, it appears that the novel AAVB1 vector is superior to AAV9 in targeting the upper airway and respiratory system, as evidenced by the enhanced ventilatory parameters during a respiratory challenge. In
addition, the improved gene delivery by AAVB1 to the tongue enhanced glycogen clearance and resulted in significant weight gain in comparison to AAV9- and PBS-treated Gaa<sup>-/-</sup> animals.

**AAV gene therapy**

AAV gene delivery offers a promising therapeutic option for monogenetic disorders such as Pompe disease, since a single intravenous administration has the potential to deliver the absent or deficient gene to the affected cells and results in an endogenous production of the missing GAA enzyme. The current FDA-approved therapy, ERT, requires weekly or biweekly infusions of rhGAA for the lifetime of the patient, as well as prolonged immune suppression. Although ERT leads to prolonged survival, many patients continue to have respiratory insufficiency secondary to untreated CNS involvement and ineffective enzyme uptake into skeletal muscle fibers. Systemic delivery of AAV9 vectors is an effective approach to treat both the muscle and CNS pathology, as these vectors can cross the blood–brain barrier. In several animal studies, AAV9 has successfully transduced muscle and motor neurons when injected directly into the muscle or intra-pleural cavity. In addition, systemic administration of AAV1-hGAA in neonatal Gaa<sup>-/-</sup> mice resulted in significant clearance of glycogen and enhanced respiratory measures. Furthermore, a previous study of AAV9-hGAA in adult Gaa<sup>-/-</sup> mice was found to be superior to ERT in respiratory outcome measures, which was attributed to effective motor neuron transduction. Although glycogen accumulation remained in the myocardium and diaphragm of AAV9-hGAA-injected animals, that study underscored the importance of treating the CNS. In this study, 3-month-old adult Gaa<sup>-/-</sup> mice were chosen to assess if gene therapy can clear the glycogen accumulation that is already present. To date, AAV gene therapy studies in adult mice with Pompe disease have had a limited transduction of respiratory and cardiac muscles, and motor neurons. Here, robust expression, enzyme activity, and clearance of glycogen in the myocardium and diaphragm are shown with both AAVB1 and AAV9.

**Strength and survival**

Higher mortality rates of Gaa<sup>-/-</sup> mice have been observed by others. Both AAVB1 and AAV9 resulted in significant cardiac transduction and increased survival. The desmin promotor has robust cardiac activity, which may account for the almost complete clearance of glycogen seen in both treatment groups. Transduction in the gastrocnemius muscle was similar to WT animals, and glycogen clearance occurred in many fibers. Nonetheless, the animals continued to have abnormal grip strength and a decreased ability to hang on to the wire net in the inverted screen test. Recently, Lee et al. used AAVN to target CNS disease and found that intracerebroventricular injections of AAVN-GAA resulted in endogenous GAA production in brain and spinal cord neurons. However, glycogen accumulation persisted in some of the motor neurons, and their muscle strength did not improve significantly. Similarly, moderate transduction efficiency in motor neurons and muscle was found in the present study, but this does not seem to be enough to reverse weakness and improve strength. In the future, exploration of a promoter that has greater expression in skeletal muscle, as well as cardiac and motor neuron activity, may enhance strength as well as prolong survival in Pompe disease.

**Tongue and respiratory pathology**

AAVB1 transduced the tongue and the hypoglossal motor neurons innervating the genioglossal muscle. The tongue plays a vital role in adequate nutrition and in maintaining upper airway patency. Contraction of the extrinsic tongue muscles dilates and stiffens the pharyngeal lumen, thereby minimizing airway narrowing and/or collapse in the face of negative inspiratory pressures. Tongue pathology is a significant cause of morbidity in IOPD and is an important therapeutic target. Tongue weakness leads to a weak suck, feeding problems early in life, and an increased risk for aspiration pneumonia. In LOPD, tongue weakness has also been documented, even in asymptomatic patients with a family history of Pompe disease. These clinical findings underscore the importance of ensuring that novel therapies adequately treat the tongue pathology. In the Pompe mouse, a direct intra-lingual injection of AAV9-hGAA can treat the tongue and the hypoglossal motor neurons, resulting in enhanced weight gain and sustained correction. Similar improvements in weight gain occurred with systemic AAVB1 therapy, presumably due to efficient vector transduction of the tongue, with subsequent clearance of glycogen in the tongue, which led to increased muscle function, allowing a greater food intake. In contrast, systemic AAV9 did not transduce the tongue as well as AAVB1. Interestingly, the Ti in AAVB1-treated animals was significantly lower than in PBS-treated mice, and it was comparable to WT animals. The decrease in Ti is a reflection of decreased upper airway resistance and improved airflow past the tongue.
Pompe disease often results in respiratory insufficiency in both IOPD and LOPD patients. Respiratory insufficiency occurs secondary to upper airway and respiratory muscle and CNS pathology. Several studies exploring respiratory directed therapy with AAV were very encouraging and resulted in a clinical trial of diaphragmatic injections of AAV1-hGAA administered into the diaphragms of children with Pompe disease. This clinical trial proved that AAV gene therapy was safe and resulted in a significant increase in the duration of time these children tolerated unassisted ventilation. In the current study, both AAVB1 and AAV9 transduced the diaphragm and the cervical ventral motor neurons where the phrenic motor neurons are located. However, during a respiratory challenge, the AAVB1 treatment group outperformed the AAV9 group, with a tidal volume and peak inspiratory flow comparable to WT animals. Both of these respiratory parameters are a reflection of inspiratory muscle strength as well as increased upper airway patency, which allows for more efficient air entry, resulting in increased tidal volume and inspiratory flow.

CONCLUSIONS

In conclusion, AAVB1-hGAA is a novel vector with muscle and CNS transduction that is superior to AAV9-hGAA in an adult Pompe disease mouse model. In this model, AAVB1-hGAA robustly decreased glycogen in the tongue and resulted in improved weight gain. In addition, Gaa−/− animals treated with AAVB1-hGAA had improved inspiratory muscle strength 6 months after treatment. Finally, although the immunological impact of both vectors in this model were not studied, when Choudhury et al. assessed the immunogenic potential of AAVB1, they found that AAVB1 was less prone to neutralization by a pooled human immunoglobulin G. This is important to consider when adult, late onset patients with Pompe disease are being targeting where pre-existing immunity is a concern. Thus, based on all these factors, AAVB1 offers a potential therapeutic option for Pompe disease.

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AUTHOR DISCLOSURE

No competing financial interests exist.

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