Molecular Bases of Disease: Glycosylation-independent Lysosomal Targeting of Acid α-Glucosidase Enhances Muscle Glycogen Clearance in Pompe Mice

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Glycosylation-independent Lysosomal Targeting of Acid α-Glucosidase Enhances Muscle Glycogen Clearance in Pompe Mice*

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Background: Acid α-glucosidase, an enzyme replacement therapy for Pompe disease, is poorly targeted to lysosomes when relying on phosphomannose residues.

Results: Fusing IGF-II to acid α-glucosidase resulted in more efficient uptake and glycogen clearance from muscle of Pompe mice.

Conclusion: Enhanced binding to the cation-independent mannose 6-phosphate receptor (CI-MPR) enabled improved glycogen clearance in Pompe mice.

Significance: BMN 701 is now being tested for Pompe disease in human clinical studies.

We have used a peptide-based targeting system to improve lysosomal delivery of acid α-glucosidase (GAA), the enzyme deficient in patients with Pompe disease. Human GAA was fused to the glycosylation-independent lysosomal targeting (GILT) tag, which contains a portion of insulin-like growth factor II, to create an active, chimeric enzyme with high affinity for the cation-independent mannose 6-phosphate receptor. GILT-tagged GAA was taken up by L6 myoblasts about 25-fold more efficiently than was recombinant human GAA (rhGAA). Once delivered to the lysosome, the mature form of GILT-tagged GAA was indistinguishable from rhGAA and persisted with a half-life indistinguishable from rhGAA. GILT-tagged GAA was significantly more effective than rhGAA in clearing glycogen from numerous skeletal muscle tissues in the Pompe mouse model. The GILT-tagged GAA enzyme may provide an improved enzyme replacement therapy for Pompe disease patients.

Pompe disease is a lysosomal storage disorder caused by insufficient lysosomal acid α-glucosidase (GAA) activity (1–3). GAA hydrolyzes the α-1,4 and α-1,6 linkages in branched glycogen molecules (4). GAA deficiency results in the accumulation of excess glycogen primarily in the lysosomes of skeletal and cardiac muscle, leading to progressive muscle weakness and cardiorespiratory failure. There is a broad spectrum in the severity and age of onset of Pompe disease that roughly correlates with the residual amount of GAA activity present in patient tissues (5). The severe infant-onset Pompe disease is characterized by hypertrophic cardiomyopathy, respiratory insufficiency, and general muscle weakness with death usually resulting by 1 year of age when untreated (6). Late-onset Pompe disease is characterized by progressive muscle weakening with respiratory failure the most common cause of death (7).

Currently, enzyme replacement therapy using alglucosidase alfa, recombinant human GAA (rhGAA) produced in Chinese hamster ovary (CHO) cells is an approved treatment for patients with Pompe disease. Generally, the clinical experience with alglucosidase alfa suggests that enzyme replacement therapy is more effective in addressing the cardiomyopathy in infant-onset patients and less effective in treating the cellular pathology of skeletal muscle in infant- and late-onset patients (6, 8–11).

In the Pompe mouse knock-out model (12), glycogen is also cleared from heart tissue more readily than from skeletal muscle (13–15). In fact, rhGAA doses of 100 mg/kg weekly were required to achieve maximal clearance of glycogen from skeletal muscle, and treatment of Pompe mice with multiple weekly doses of 20 mg/kg rhGAA cleared only between 0 and 40% of glycogen from quadriceps.

The marketed dose of alglucosidase alfa (20 mg/kg biweekly) is relatively high compared with the dose of approved therapeutics for other lysosomal storage diseases (biweekly doses between 0.5 and 1 mg/kg for Gaucher, Fabry, and mucopolysaccharidosis I), suggesting that delivery of rhGAA to the lysosome of skeletal muscle is of limited efficiency. In part, this is because of the intrinsic properties of skeletal muscle, such as the low...
abundance of the cation-independent mannose 6-phosphate receptor (CI-MPR), altered lysosomal function leading to increased autophagy particularly in type II muscle fibers (19, 20), and the relatively low blood flow to skeletal muscle tissue compared with liver, heart, and other tissues. However, the problem is exacerbated by the low affinity of CHO-produced rhGAA for the CI-MPR, which results in diminished lysosomal targeting from the extracellular pathway (3, 21, 22).

rhGAA is delivered to the lysosome by receptor-mediated endocytosis after binding of bis-mannose 6-phosphorylated glycan (bis-M6P-glycan) on the rhGAA protein surface to the CI-MPR. rhGAA has only about 0.9–1.2 mol M6P/mol enzyme, and only a small fraction of rhGAA oligosaccharides have bis-M6P (14, 15). As oligosaccharides with only a single M6P bind to the CI-MPR with a 1000-fold lower affinity (24), rhGAA preparations containing less than 0.9 mol M6P/mol protein are expected to have decreased targeting to the lysosomes.

We have used a peptide-based, glycosylation-independent lysosomal targeting (GILT) approach to create an rhGAA molecule containing a high affinity ligand for the CI-MPR. Previously, fusion of a portion of human IGF-II to the human lysosomal enzyme β-glucuronidase, deficient in mucopolysaccharidosis VII (MPS VII), was shown to promote efficient uptake into cultured MPS VII fibroblasts and delivery of active enzyme to a wide range of cell types in MPS VII mice through binding of the IGF-II-derived GILT tag to the CI-MPR (26).

We fused the GILT tag to the N terminus of rhGAA to create a GAA enzyme (BMN 701) with high affinity for the CI-MPR receptor. BMN 701 has biochemical properties similar to those of rhGAA but is delivered more effectively to the lysosomes of L6 myoblasts in vitro. Administration of GILT-tagged BMN 701 to Pompe model mice greatly increases the clearance of accumulated glycogen in muscle tissue compared with equivalent doses of rhGAA. Use of the GILT peptide tag addresses the relatively ineffective targeting of rhGAA caused by poor M6P phosphorylation, and the BMN 701 molecule is now being tested in a phase I-II clinical study to determine whether its increased efficiency of targeting will improve clinical outcomes.

EXPERIMENTAL PROCEDURES

Plasmids—DNA cassette 635 encoding complete human GAA amino acids 1–952 was derived from IMAGE clone 4374238 (Open Biosystems) using PCR primers GAA13 (5′-gtcttgaCTAACACGCTGACCCGCTTCG-3′) and GAA27 (5′-gtcgtgaCTAACACCAGCTGACCCGCTTCG-3′). This cassette was digested with EcoRI and XbaI, blunted by treatment with Klenow DNA polymerase, and then ligated into the Klenow-treated HindIII site of expression vector pCEP4 (Invitrogen) to create plasmid p635.

DNA cassette 701 was identical to cassette 635 except for the following GILT tag-containing N-terminal sequence that was joined upstream of GAA amino acid Ala-70: gaattcCACCCATCCCTTTTGCAGGTGTGCGATGCATTGT-TCACCTTTTATGGCCTCGTCCTGGTCTGCGATTCATGCTGCTGTGTTTCCGCAGCCCTCGTCACTTCCTGGGAGCTGACGAGAAAC-CTACACCATCATCGCTGGTGCTTCTC-3′/H11032.

Plasmid 701 cassette in CHO cells was generated with the GPEx™ retrovector expression system as described (27). The 5′-end of DNA cassette 1288 (gaattcACACCAATGGGAATCTGACTGCTGGTGCTTCTC-ACCTCTTGTGGGCTCGTCACTTGGCTGCGATTCATGCTGCTGTCGCGCAGCCCTCGTCACTTCCTGGGAGCTGACGAGAAAC-CTACACCATCATCGCTGGTGCTTCTC-3′/H11032) and an increase in glycogen clearance from Pompe mouse muscle tissue (14, 25).

Delivery of rhGAA can be improved by providing high affinity ligands for the CI-MPR. One approach used was the conjugation of bis-M6P-containing oligosaccharides to rhGAA. This neo-rhGAA displayed an increase in cellular uptake in vitro and in vivo and an increase in glycogen clearance from Pompe mouse muscle tissue (14, 25).

Tissue Culture—For many experiments, GILT-tagged GAA and rhGAA were purified from culture supernatants of transiently transfected HEK293 cells. For transient transfections, plasmids p701 and p635 were transfected into suspension FreeStyle 293-F cells as described by the manufacturer (Invitrogen). Briefly, cells were grown in Opti-MEM I medium (Invitrogen) in polycarbonate shaker flasks on an orbital shaker at 37 °C 80% CO2. Cells were then adjusted to a concentration of 1×10^6 cells/ml and then transfected with a 1:1:1 ratio of ml cells:μg DNA:μl 293Fecin. Cells were harvested 5–10 days post-transfection and centrifuged at 5,000 × g for 20 min. Culture supernatants were filtered through a 0.8/0.2-μm AcroPak 500 capsule ( Pall) and concentrated with a tangential flow device with a 30,000 MWCO PLTK filter (Millipore). Sodium acetate, pH 4.6, was added to a final concentration of 0.1 M, and concentrated supernatants were quick-frozen in liquid nitrogen and stored at −80 °C.

A stable cell line suitable for large scale production of the GILT-tagged 701 cassette in CHO cells was generated with the GPE™ retrovector expression system as described (27). The resulting BMN 701 CHO cell line was grown in 5 liters of PF-CHO LS medium (HyClone) in a 10-liter Wave bioreactor at 37 °C in 5% CO2. Cultures were supplemented with R15.4
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(HyClone). On day 5 cultures were shifted to 31 °C and then harvested on day 11 and frozen at −80 °C.

L6 rat myoblasts (ATCC, CRL-1458) and C2C12 mouse myoblasts (ATCC, CRL-1772) were grown in DMEM (Invitrogen) supplemented with 10% fetal bovine serum (Invitrogen) in 75-cm² tissue culture flasks (Corning) at 37 °C and 5% CO₂.

CI-MPR Biotinylation—Supernatant from FreeStyle 293-F cells transfected with plasmids p1288 and p1355 were applied to a 1-ml His-GraviTrap column (GE Healthcare) as directed by the manufacturer for purification of the His₆-tagged receptor domain proteins. Eluted proteins were concentrated, exchanged into 10 mM Tris, pH 8, and 25 mM NaCl buffer, and then biotinylated with BirA enzyme as described by the manufacturer (Avidity) in reactions that contained 70 µg of receptor in 205 µl, 10 mM Tris, pH 8, 25 mM NaCl buffer, 25 µl of BiomixA, 25 µl of BiomixB, and 4 µl of BirA enzyme. BirA enzyme treatment was performed at 30 °C for 1.5 h. The reactions were then diluted 20-fold into His GraviTrap binding buffer (GE Healthcare) and reapplied to the His GraviTrap column for removal of BirA enzyme and free biotin. Biotinylated CI-MPR constructs were stored at 4 °C.

Protein Purification—Cell culture supernatants adjusted to pH 4.6 with sodium acetate were thawed from storage at −80 °C, and ammonium sulfate was added to 0.75 M. The material was centrifuged to remove precipitation and filtered with a 0.8/0.2-µm AcroPak 500 capsule (Pall). The filtered material was loaded onto a phenyl-Sephase 6 Fast Flow (low sub) (GE Healthcare) column prepared with HIC load buffer (50 mM sodium acetate, pH 4.6, 0.75 M ammonium sulfate). The column was washed with 10 column volumes of HIC wash buffer (50 mM sodium acetate, pH 5.3, 0.75 M ammonium sulfate) and eluted with 5 column volumes of HIC elution buffer (50 mM sodium acetate, pH 5.3, 20 mM ammonium sulfate). Pooled fractions were dialyzed extensively into QXL load buffer (20 mM histidine, pH 6.5, 50 mM NaCl) and then loaded onto a Q Sepharose XL column (GE Healthcare). The column was washed with 10 column volumes of QXL load buffer and eluted with 10 column volumes of QXL elution buffer (20 mM histidine, pH 6.5, 150 mM NaCl). Purified samples were concentrated by ultrafiltration, and buffer was exchanged into phosphate-buffered saline, pH 6.2. Samples were snap-frozen in liquid nitrogen and stored at −80 °C. Alg glucosidase alfa was recovered from excess material of the reconstituted commercial product used in patients.

Enzyme Assays—Assay substrates and standards were obtained from Sigma. For GAA para-nitrophenol (PNP) assays, enzyme was incubated in 50-µl reactions containing 100 mM sodium acetate, pH 4.2, with 10 mM PNP α-glucoside substrate. Reactions were incubated at 37 °C for 20 min and stopped with 300 µl of 100 mM sodium carbonate. Absorbance at 405 nm was read in 96-well microtiter plates and compared with standard curves derived from p-nitrophenol. 1 GAA PNP unit is defined as nmol PNP hydrolyzed/ml enzyme/h. Specific activities were determined by dividing GAA PNP units by the protein concentration as determined by the Bradford assay (Bio-Rad). Saturation curves were generated by incubating 0.5 µg of GAA protein in PNP assays with increasing amounts of PNP α-glucoside ranging from 0 to 20 mM substrate. Plots of product formation versus substrate concentration were fit to the Michaelis-Menten equation with KaleidaGraph software (Synergy Software) to determine Kₘ values.

GAA was also assayed using 4-methylumbelliferyl α-D-glucosidase substrate. The enzyme was incubated in 20-µl reactions containing 123 mM sodium acetate, pH 4.0, with 10 mM 4-methylumbelliferyl α-D-glucosidase substrate. Reactions were incubated at 37 °C for 1 h and stopped with 200 µl of buffer containing 267 mM sodium carbonate and 427 mM glycine, pH 10.7. Fluorescence was read with 355 nm excitation and 460 nm filters in 96-well microtiter plates and compared with standard curves derived from 4-methylumbelliferone. One GAA 4-methylumbelliferyl unit is defined as nmol 4-methylumbellifereone hydrolyzed/ml/h.

Uptake Assays—L6 myoblasts were seeded into 24-well tissue culture plates (Corning), grown for 24 h to ~90% confluency. Cells were washed once with uptake medium (DMEM, high glucose, supplemented with 0.5 g/liter bovine serum albumin (Sigma), 4 mM L-glutamine (Invitrogen), and 20 mM HEPEs, pH 7.2), and then incubated at 37 °C and 5% CO₂ in uptake medium containing purified rhGAA or BMN 701. Some wells also contained either 5 mM M6P (Calbiochem) or 2.4 mM IGF-II (Cell Sciences) as an inhibitor. After 18 h, cells were washed four times with DPBS, pH 7.4 (0.133 g/liter CaCl₂, 2H₂O, 0.1 g/liter MgCl₂·6H₂O, 0.2 g/liter KCl, 0.2 g/liter KH₂PO₄, 8.0 g/liter NaCl, 1.15 g/liter Na₂HPO₄), and then lysed with the addition of 200 µl/well Cell.Lytic M reagent (Sigma) and 20 min of shaking at room temperature. Lysate debris was removed by centrifugation at 2,000 × g for 5 min. Lysate GAA activity of duplicate wells was measured with the GAA 4-methylumbelliferyl assay and normalized to lysate protein concentration as determined using the bicinchoninic acid protein assay (Pierce).

SDS-PAGE, Western Blot Analysis, and Antibodies—Protein samples were separated on Criterion XT 4–12% BisTris reducing SDS-PAGE (Bio-Rad) at 200 V for 50 min as described by the manufacturer. Silver staining was performed as described (29). For Western blots, samples were transferred to Hybond-P membranes (GE Healthcare) and probed using the ECL system (GE Healthcare) as described by the manufacturer. Dilution of primary anti-GAA mouse monoclonal 3A6-1F2 antibody was 1:100,000. Rabbit polyclonal anti-IGF2 (Cell Sciences, PA0382) and anti-GAA PAb2443 were diluted 1:1000 and 1:200, respectively. Horseradish peroxidase-conjugated secondary anti-mouse IgG and anti-rabbit IgG antibodies (BioChain) were diluted 1:5,000. Molecular weight markers were from Bio-Rad. Anti-GAA monoclonal antibody 3A6-1F12 was generated using rhGAA as antigen, and antibody used in this study was purified from ascites fluid by QED Bioscience. Polyclonal antibody 2443 was produced in rabbits using rhGAA excised from an SDS-polyacrylamide gel as antigen.

Immunofluorescence—C2C12 mouse myoblasts were grown on polylysine-coated slides (BD Biosciences) and incubated for 18 h in the presence or absence of 100 nM BMN 701 at 37 °C in 5% CO₂. Cells were incubated in growth medium for 1 h and then washed four times with DPBS before fixing with methanol at room temperature for 1 h. The following incubations were all at room temperature, each separated by three washes in DPBS.
Incubations were for 1 h unless noted. Slides were permeabilized with 0.1% Triton X-100 for 15 min and then blocked with blocking buffer (10% heat-inactivated horse serum (Invitrogen) in DPBS). Slides were incubated with primary mouse monoclonal anti-GAA antibody 3A6-1F2 (1:5000 in blocking buffer) and then with secondary rabbit anti-mouse IgG AF594-conjugated antibody (Invitrogen A11032, 1:200 in blocking buffer). A FITC-conjugated rat anti-mouse LAMP-1 (BD Pharamingen 553793, 1:50 in blocking buffer) was then incubated. Slides were mounted with DAPI-containing mounting solution (Invitrogen) and viewed with a Nikon Eclipse 80i microscope equipped with fluorescein isothiocyanate, Texas Red, and DAPI filters (Chroma Technology). Images were captured with a photometric Cascade camera controlled by MetaMorph software (Universal Imaging). Images were merged using Photoshop software (Adobe).

Surface Plasmon Resonance Analysis—All surface plasmon resonance measurements were performed at 25 °C using a BIAcore 3000 instrument. SA sensor chips and surfactant P20 were obtained from GE Healthcare. All buffers were filtered using Nalgene filtration units (0.2 μm), equilibrated to room temperature, and degassed immediately prior to use. Protein samples were centrifuged (15 min at 13,000 × g) to remove any particulates that might be present in the sample. Purified biotinylated wild-type (1288) or mutant (1355; containing a single amino acid substitution that effectively decreases the affinity of the receptor for IGF-II) CI-MPR constructs were immobilized on separate surfaces on a BIAcore streptavidin SA chip to a final

for glycogen level analysis. Of the 40 initial mice (5 animals/group), 8 died during the experiment: 2 control animals, 2 animals from each of the low dose groups (rhGAA and BMN 701, 5 mg/kg), and 1 animal from each high dose group (rhGAA and BMN 701 20 mg/kg). The animals were reared in groups with no evidence of chronic illness. Animals died evenly across all groups, so the animal deaths are not thought to be compound related.

Tissues were homogenized using a rotor/stator homogenizer and then measured for glycogen content in a blinded format using the Amplex Red glucose assay kit (Invitrogen) essentially as described (14). For each tissue site, an analysis of variance was performed to obtain a pooled estimate of variance, and two-sided t tests within the analysis of variance model were performed to compare each pair of the five treatments, for a total of 10 comparisons. p values less than 0.05 are considered nominally statistically significant. GAA activity levels were measured in duplicate from the homogenized tissues, and the averages are reported as units/mg protein in Table 2.

RESULTS

GILT-GAA Has Enzymatic Activity Comparable with Untagged GAA—Fig. 1 shows SDS-PAGE of rhGAA and GILT-tagged-GAA (BMN 701) purified from culture supernatants of transiently transfected suspension HEK293 cells. SDS-PAGE analysis shows that the purified rhGAA preparation is predominantly the 110,000-kDa full-length precursor and contains detectable amounts of the 95- and 75-kDa processed species. HEK293-produced BMN 701, which contains the GILT tag fused to position 70 of GAA, migrates with A slightly higher molecular mass. Western blot analysis with anti-GAA antibodies (Fig. 1B) and an anti-IGF2 tag antibody (Fig. 1C) confirms the identity of the GAA polypeptide in both samples and the presence of the GILT tag in purified BMN 701.

The enzymatic activity of the two compounds was measured using a synthetic substrate, PNP α-glucoside. The specific activities of rhGAA and BMN 701 were 143,000 and 140,000 units/mg, respectively. Specific activities for CHO-produced BMN 701 ranged from 137,000 to 150,000 units/mg. Saturation
curves yielded similar $K_m$ values of 5.8 and 5.9 mM for rhGAA and BMN 701, respectively.

The GILT Tag Mediates Uptake of BMN 701 into Cultured Myoblasts—Fig. 2 shows the ability of excess M6P or IGF-II to inhibit uptake of BMN 701 and rhGAA into rat L6 myoblasts. Cellular uptake of GILT-tagged BMN 701 is not inhibited by excess M6P but is inhibited completely by excess IGF-II. These data are consistent with the prediction that uptake of BMN 701 would occur by binding of the GILT tag to the IGF-II binding site of the CI-MPR. Uptake of rhGAA is inhibited completely by M6P, indicating that its internalization relies solely on M6P recognition by the M6P binding sites on the CI-MPR. Excess IGF-II partially competes for uptake of rhGAA. This phenomenon has been reported previously for other M6P-bearing lysosomal enzymes and has been attributed to steric hindrance of the M6P sites when IGF-II engages its cognate binding site on the receptor (31).

The specificity of the receptor-mediated uptake of the GILT-tagged BMN 701 protein was confirmed by uptake inhibition studies using two protein competitors derived from the CI-MPR (Fig. 3). Receptor construct 1288 contains domains 10–13 of the CI-MPR, sufficient to bind IGF-II but not M6P (32, 33). Receptor construct 1355 is identical to 1288 except for a single amino acid substitution that decreases its affinity for IGF-II (34). Neither receptor fragment inhibits uptake of rhGAA (Fig. 3A), indicating that rhGAA does not bind to the IGF-II binding domain of the CI-MPR. In contrast, construct 1288 is a potent inhibitor of BMN 701 uptake, and 1355 is a poor inhibitor (Fig. 3B). These data prove that uptake of BMN 701 occurs through interaction of the GILT tag with the IGF-II binding domain of the CI-MPR.

Intracellular Processing of GILT-tagged and Untagged GAA Is Similar—The kinetics of GAA enzyme activity decay after uptake of rhGAA and BMN 701 into L6 myoblasts were determined over a 14-day period (Fig. 4). The proteins have comparable half-lives of 8.3 and 9.0 days for rhGAA and BMN 701, respectively.

Once internalized, the 110-kDa precursor form of rhGAA is processed sequentially in the lysosome to an intermediate 95-kDa form, then to the mature 76- and 70-kDa forms. Smaller
mouse myoblasts were cultured in the presence and absence of GILT-tagged BMN 701. Fluorescently tagged antibodies against human GAA and mouse lysosomal marker LAMP1 (36) were then used to probe the cells (Fig. 6). Anti-GAA antibody detects internalized protein in cells cultured with BMN 701 but not in cells grown in the absence of BMN 701. The staining pattern of the anti-GAA antibody in individual cells grown with BMN 701 is indistinguishable from the staining pattern of anti-LAMP1, indicating that GILT-tagged BMN 701 is internalized and trafficked to the lysosome.

**GILT-tagged GAA Has Enhanced Delivery to Myoblasts Compared with Alglucosidase Alfa—** *In vitro* rat L6 myoblast cell uptake assays were performed with increasing amounts of BMN 701 to generate uptake saturation curves (Fig. 7). For comparison, uptake saturation curves were generated with untagged rhGAA, which was also used in animal testing, and with the currently approved Pompe enzyme replacement therapy, alglucosidase alfa, manufactured by Genzyme Corp., which was unavailable for animal testing. The cell uptake kinetics of HEK293-produced rhGAA and alglucosidase alfa are indistinguishable, with \( K_{\text{uptake}} \) values of 141 and 147 nM, respectively. GILT-tagged BMN 701, however, shows a 26-fold enhancement in cellular delivery compared with alglucosidase alfa or rhGAA, with a \( K_{\text{uptake}} \) value of 5.4 nM. Additionally, the BMN 701 uptake saturated reproducibly at a level \( \approx \) 2-fold greater than either of the untagged enzymes (1100 units/mg for BMN 701 compared with 619 and 516 units/mg for alglucosidase alfa and rhGAA, respectively).

**GILT-tagged GAA Has High Affinity for the CI-MPR—** To gauge the affinity of GILT-tagged BMN 701 for the IGF-II binding site on the CI-MPR, we compared its binding affinity with that of full-length recombinant IGF-II using surface plasmon resonance (Fig. 8). Association with the IGF-II binding domain (domains 10–13) of receptor fragment 1288 was measured using receptor 1355 (I1572T mutant) as a negative binding control. The \( K_d \) values for the interaction of BMN 701 and IGF-II were determined by both a 1:1 Langmuir fit of the association and dissociation rate constants and by steady state kinetics (Table 1 and Fig. 8). The \( K_d \) determined for the interaction of IGF-II with the domain 10–13 construct is 2.5 nM by Langmuir and 8.8 nM by steady state kinetics. The \( K_d \) determined for interaction of BMN 701 with the domain 10–13 construct is 16 nM by Langmuir and 21 nM by steady state kinetics, within 6.4- and 2.4-fold, respectively, of the \( K_d \) values determined for IGF-II. Linnell et al. (33) reported that IGF-II binds to a domain 10–13 construct with a \( K_d \) 8.6-fold greater than for binding to domains 1–15 of the receptor, indicating that other domains of the receptor stabilize the interaction. If this ratio held for BMN 701, then the \( K_d \) for binding to the intact receptor would be about 2 nM, a value that is in good agreement with the \( K_{\text{uptake}} \) determined in uptake studies. Other literature values of the \( K_d \) for IGF-II binding the complete CI-MPR range from 0.2 to 16 nM (37–39). Considering the tighter binding of IGF-II for the intact receptor, the determined \( K_d \) value for IGF-II binding to the domain 10–13 construct is consistent with the lower end of this range. Binding of rhGAA to the domain 10–13 construct was not detected (data not shown).
GILT-tagged GAA Enhances Clearance of Glycogen from Mouse Muscle Tissue Compared with rhGAA—The abilities of BMN 701 and rhGAA to reverse the storage pathology in Pompe mice was tested by administering the compounds to mice in a 4-week enzyme replacement protocol. The uptake kinetics of the HEK-produced rhGAA used in this experiment were evaluated as described above (Fig. 7).

Mice received four injections of BMN 701 or rhGAA at doses of either 5 or 20 mg/kg. The reduction in glycogen storage was greater in both dosing groups receiving BMN 701 compared with those receiving rhGAA (Fig. 9). In all tissues, treatment with the 20 mg/kg dose of BMN 701 resulted in statistically significant ($p < 0.05$) lower levels of glycogen than either the 20 or 5 mg/kg dose of rhGAA. Except for in the lingual musculature, treatment with 5 mg/kg BMN 701 resulted in significantly ($p < 0.05$) lower levels of glycogen than the 5 mg/kg dose of rhGAA. In the tibialis anterior, extensor digitorum longus, gastrocnemius, heart, and quadriceps, treatment with the 5 mg/kg dose of BMN 701 resulted in significantly ($p < 0.05$) lower levels of glycogen than treatment with 20 mg/kg rhGAA. No correlation was found between glycogen clearance and tissue GAA activity (Table 2). This is presumably because of the accumulation of significant quantities of GAA enzyme in capillary endothelial cells in muscle tissue (15).

**DISCUSSION**

The rationale for enzyme replacement therapy for Pompe disease is that the underlying molecular cause of Pompe disease, accumulation of excess glycogen in the lysosomes of car-
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TABLE 1
Surface plasmon resonance analysis

<table>
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<tr>
<th>Ligand</th>
<th>$k_a$</th>
<th>$k_d$</th>
<th>$K_d$</th>
<th>$R_{max}$</th>
<th>Steady-state kinetics</th>
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<tr>
<td></td>
<td>1/ms</td>
<td>1/s</td>
<td>nM</td>
<td>RU</td>
<td>$K_d$</td>
</tr>
<tr>
<td>BMN 701</td>
<td>$3.6 \times 10^3 \pm 1.9 \times 10^2$</td>
<td>$5.8 \times 10^{-2} \pm 1.4 \times 10^{-2}$</td>
<td>16</td>
<td>146</td>
<td>21 $\pm$ 1.9</td>
</tr>
<tr>
<td>IGF-II</td>
<td>$1.1 \times 10^6 \pm 8.8 \times 10^3$</td>
<td>$3.4 \times 10^{-3} \pm 2.8 \times 10^{-3}$</td>
<td>2.5</td>
<td>16.1</td>
<td>8.8 $\pm$ 1.2</td>
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BMN 701 is expressed containing the IGF-II signal sequence, and rhGAA is expressed containing the native GAA signal sequence. Even though native GAA is poorly phosphorylated, replacement of the native signal sequence with that from IGF-II is likely to change the post-translational glycosylation. In fact substitution of the IGF-II signal peptide for the endogenous GAA signal peptide results in a GAA molecule with diminished M6P-dependent uptake, possibly because of alterations in the kinetics of GAA trafficking (data not shown). The native GAA signal peptide is reported to be inefficiently cleaved (40), which could extend its residence time in the Golgi, where M6P addition occurs.

Finally, BMN 701 is significantly more potent than untagged rhGAA in muscle tissue glycogen clearance in the Pompe mouse model (Fig. 9). In most muscle tissues examined, BMN 701 doses of 5 mg/kg led to greater clearance of glycogen than untagged rhGAA doses of 20 mg/kg. BMN 701 appears most effective in clearing glycogen from the heart, soleus, tibialis anterior, extensor digitorum longus, gastrocnemius, and quadriiceps. A sufficient amount of commercially available rhGAA (alglucosidase alfa) could not be obtained for this animal experiment, and thus, despite the fact that the uptake kinetics in rat L6 myoblasts of the rhGAA used in this study were virtually indistinguishable from those of alglucosidase alfa (Fig. 7), the possibility of other differences between the two untagged rhGAs, such as differences in the sialic acid content of the glycan, limited the ability to draw conclusions about the relative performance of the two rhGAs in animals.

Current enzyme replacement therapies for the treatment of lysosomal storage diseases rely on the binding of rhGAA to the CI-MPR via M6P moieties on the glycosylated rhGAA protein. The work presented here shows an alternative route for engagement of the receptor via an IGF-II-derived GILT tag fused to the rhGAA molecule. This alternative may hold several advantages over current enzyme replacement therapy for Pompe disease. For example, use of the GILT targeting strategy is compatible with non-mammalian expression systems that do not produce M6P-containing oligosaccharides on proteins. Also, because the glycosylation state of proteins can be variable and influenced by cell culture conditions, large scale production of enzymes that relies on specific glycosylation for their biological activity, such as M6P-dependent targeting, can face technical challenges. This appears to have been the case with alglucosidase alfa, where differences in pharmacokinetics and biodistribution between lots produced at the 160-liter scale and the 2000-liter scale were attributed to differences in glycosylation by the FDA, leading to the filing of a new biological license application for the 2000-liter–produced rhGAA (41).

Another potential advantage of BMN 701 relates to the observation that uptake of BMN 701 into rat L6 myoblasts sat-
Figure 9. Glycogen clearance in Pompe mice. Glycogen levels were measured in various muscle tissues following four injections with CHO-produced BMN 701 or rhGAA. See “Experimental Procedures” for details.
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