Patients with α-dystroglycanopathies, a subgroup of rare congenital muscular dystrophies, present with a spectrum of clinical manifestations that includes muscular dystrophy as well as CNS and ocular abnormalities. Although patients with α-dystroglycanopathies are genetically heterogeneous, they share a common defect of aberrant post-translational glycosylation modification of the dystroglycan alpha-subunit, which renders it defective in binding to several extracellular ligands such as laminin-211 in skeletal muscles, agrin in neuromuscular junctions, neurexin in the CNS, and pikachurin in the eye, leading to various symptoms. The genetic heterogeneity associated with the development of α-dystroglycanopathies poses significant challenges to developing a generalized treatment to address the spectrum of genetic defects. Here, we propose the development of a bispecific antibody (biAb) that functions as a surrogate molecular linker to reconnect laminin-211 and the dystroglycan beta-subunit to ameliorate sarcolemmal fragility, a primary pathology in patients with α-dystroglycan-related muscular dystrophies. We show that the treatment of LARGE

**INTRODUCTION**

α-dystroglycanopathy is the collective term for a clinically and genetically heterogeneous group of muscular dystrophies with the biochemical hallmark of decreased or absent specific O-linked glycosylation on α-dystroglycan (αDG) due to genetic mutations in one of 18 genes identified thus far. The hypoglycosylation of specific O-linked glycan on αDG results in diminished binding to its ligands in the extracellular matrix, including laminin-211, agrin and perlecan in muscles, neurexin in the brain, and pikachurin in the eye. These aberrations lead to clinical manifestations in the muscles, the CNS, and the eye. Patients with severe α-dystroglycanopathy subtypes, such as Walker-Warburg syndrome (WWS), muscle-eye-brain disease (MEB), and Fukuyama-type congenital muscular dystrophy (FCMD), are typically diagnosed at birth or within a few months of life. These patients develop generalized hypotonia, developmental delays, mental retardation due to brain structural malformations, and eye abnormalities. In contrast, patients with limb-girdle muscular dystrophy (LGMD) typically acquire ambulation but develop muscle weakness in their early teens and become wheelchair bound in early or late adulthood. Due to the lack of a strict correlation between genotype and phenotype, these diseases were recently reclassified into four subtypes: (1) forms with congenital-onset weakness and severe structural brain abnormalities (WWS, MEB, and FCMD); (2) forms with congenital-onset weakness with brain involvement ranging from absent to moderate (congenital muscular dystrophy types 1C [MDC1C] and 1D [MDC1D]); (3) forms with later-onset limb-girdle weakness associated with mental retardation and microcephaly but without structural brain defects (LGMD type 2K [LGMD2K]); and (4) the mildest variants with no brain involvement (LGMD type 2I [LGMD2I]).

DG is a component of the dystrophin-associated glycoprotein complex (DGC), which primarily serves to link the extracellular matrix to the intracellular cytoskeleton, conferring sarcolemma stability during muscle contraction. α- and βDGs are transcribed from the same gene (DAG1) and are generated by post-translational cleavage of the gene product to produce a transmembrane segment (βDG) and an extracellular, non-covalently associated protein (zDG), which is heavily O-glycosylated. An important feature of this glycosylation, the mechanism of which was recently elucidated, is the unique O-linked glycans (matriglycan) that is modified by the
xylose-glucuronic acid repeats by LARGE\textsuperscript{15} that endow αDG with ligand-binding activity.\textsuperscript{16}

The genetic heterogeneity of α-dystroglycanopathy and the rarity of each individual subtype pose a significant challenge to developing a single treatment that would benefit all the subtypes. Both αDG (via the matriglycan) and integrin α7β1 bind to laminin-211 in the basal lamina of skeletal muscles and contribute to force production.\textsuperscript{17} However, most of the mechanical support for sarcolemma integrity appears to be provided by the binding of laminin-211 in the basal lamina to αDG on the sarcolemma. Hence, conceptually, replacing the non-functional hypoglycosylated αDG in α-dystroglycanopathy with a functional mimic to restore the linkage between laminin-211 and the sarcolemma may serve as a therapeutic (i.e., protein replacement) for this patient population. That protein replacement may be feasible was illustrated by the demonstration that local injection of purified matriglycan O-glycosylated αDG\textsuperscript{17} into the muscles of LARGE\textsuperscript{null} mice conferred a measure of protection to the sarcolemma from laser-induced damage. However, when the matriglycan-glycosylated αDG was injected systemically, very little change was noted in muscle tissues, presumably due to rapid clearance by the mannose or asialoglycoprotein receptors on the cells in the reticuloendothelial system. To overcome this obstacle, we elected to develop a bispecific antibody (biAb) engineered to bind simultaneously both laminin-211 and αDG and to serve as a surrogate linker that connects the basal lamina to the sarcolemma. Importantly, such an approach, if successful, should be applicable to treat several subsets of α-dystroglycanopathy patients. Here, we show that administering biAbs with this functionality into LARGE\textsuperscript{null} mice (a model of α-dystroglycanopathy) protected the animals from exercise-induced muscle damage and, consequently, improved muscle function.

RESULTS

Development of biAbs Targeting Laminin-211 and βDG

The dystrophin-associated glycoprotein complex links the intracellular cytoskeleton to the basal lamina in muscles.\textsuperscript{13} Figure 1A depicts the key components of the DGC that create this linkage. βDG, the transmembrane protein component of DG, is positioned centrally in the DGC. Intracellularly, βDG confers mechanical support through its association with dystrophin to connect the sarcolemma with actin filaments. Extracellularly, together with the sarcoglycans and biglycan,\textsuperscript{14,15} βDG associates with αDG to bind the basal lamina via its matriglycan with laminin-211. Because the integrity of skeletal muscle sarcolemma requires both intra- and extracellular support, a defect in any one of the DGC components frequently leads to muscular dystrophy.\textsuperscript{12,20} For example, dystrophin defects play a role in the development of Duchenne muscular dystrophy,\textsuperscript{21} and sarcoglycan defects play a role in the development of sarcoglycanopathies.\textsuperscript{22}

In αDG-related muscular dystrophies, the link between laminin-211 and αDG is disrupted due to a deficiency in O-linked matriglycan on αDG (Figure 1B). To address this loss of function, we envisioned the development of a biAb designed to concurrently bind laminin-211 and αDG and thereby act as a surrogate of the normal function of matriglycan on αDG. Since α-dystroglycanopathies may include a secondary deficiency in αDG levels, we reasoned that a biAb capable of recognizing laminin-211 and βDG rather than αDG might restore the linkage to a greater extent (Figure 1C). The spatial distance between βDG and laminin-211 was also anticipated to be a consideration for optimal DGC function. Providentially, the distance between the sarcolemma and laminin-211 is reportedly in the ~100-Å range,\textsuperscript{17} which approximates that of biAbs that can span up to 150 Å.

Laminin-211 comprises the three subunits α2, β1, and γ1; hence, its designation as laminin-211.\textsuperscript{23,24} The α2 subunit contains five laminin-globular (LG) domains at its C terminus that participate in binding to different extracellular molecules, such as sulfatides, integrin α7β1, and heparin;\textsuperscript{25} the fourth and fifth LG domains (LG4/5) bind αDG via the matriglycan.\textsuperscript{26} To minimize the potential interference of the biAb with the physiological associations between other LG domains and their respective binding partners, we elected to use the LG4/5 domains as the target antigens for the proposed biAb. Alignment of the human and mouse laminin-α2 LG4/5 protein sequences indicated that they shared a sequence identity of approximately 88% and that the identity between the human and mouse sequences of the extracellular domain (ECD) of βDG (96 amino acids) was 86%. Thus, it was conceivable that we could generate Abs that cross-reacted with both human and mouse laminin-α2 and βDG, obviating the need to develop two sets of Abs that separately recognized human and mouse targets. To this end, human and mouse laminin-α2 LG4/5 and the ECD of βDG were expressed in HEK293 cells, purified, and then used to generate hybridomas. To ensure that the Abs could recognize the native conformations of the epitopes, correct folding of LG4/5 was confirmed by binding to heparin.\textsuperscript{25} Monoclonal antibodies (mAbs) to laminin-α2 chain and βDG were identified and characterized using a variety of assays (Figures S1A–S1E) to ensure that they bound their respective targets with
the desired properties. The resulting mAbs positively stained unfixed frozen sections of human, mouse, and cynomolgus monkey muscle tissues, indicating that they recognized their respective targets in the native state (Figure S1C).

For an initial demonstration of proof of concept, the VH and VL sequences of the selected anti-laminin-211 and anti-βDG ECD mAbs were used to engineer biAbs in the tetravalent bispecific tandem immunoglobulin (Ig) (TBTI) format using mouse Fc as a backbone (Figure S2A). To ensure that consequent binding of the Ab to the sarcolemma or basal lamina would not induce activation of effector cells or the complement pathway, the effector function of the Fc was silenced by introducing δAB mutations. The biAbs were expressed in HEK293 cells, purified, and extensively characterized to verify that they retained the binding properties of the parental mAbs. We showed that the biAbs retained binding activity, albeit at a reduced affinity to laminin-211 than the parental mAb (Table S1).

Sandwich ELISA and sequential surface plasmon resonance (SPR) assays confirmed that the biAbs could engage both LG4/5 and βDG concurrently (Figures S2B and S2C). However, sequential SPR binding assays again revealed a reduction in the binding affinity of the anti-laminin-211 Ab in the TBTI biAb format with an inner VH/VL configuration (Figure S2B, arrowheads). This suggested that steric hindrance in the inner complementarity-determining regions may have impaired antigen binding, as predicted by crystal structure modeling.

Local Intramuscular Injection of the biAb Protects the Tibialis Anterior Muscle from Exercise-Induced Damage

To evaluate whether the biAb (recognizing anti-laminin-211 and βDG) exhibited the desired function in vivo, we used the LARGEmyd-3J mouse model of α-dystroglycanopathy (Jackson Laboratory; LARGEmyd-3J /GrsrJ, stock #008581). The LARGEmyd-3J mouse has a hypomorphic mutation in Large (like-glycosyltransferase) that leads to a loss in Large expression, consequent reduced O-linked IIH6-positive matriglycans on αDG, and an ability to bind laminin-211 (in a laminin-overlay assay) and thereby render the myofibers susceptible to contraction-induced damage. LARGE is the xylose-glucuronic acid dual glycosyltransferase critical for the addition of the xylose-glucuronic acid disaccharide repeats responsible for ligand binding. This reduction in glycosylation was observed in muscle lysates of LARGEmyd-3J mice when compared to wild-type mice using the glycosylation-specific IIH6 antibody (Figure 2A). Both αDG and βDG were also evident in the muscle lysates of wild-type and LARGEmyd-3J mice, with the latter showing a shift in the size of zDG reflecting a loss in glycosylation (Figure 2A). Phenotypically, this mouse model resembles a severe form of LGMD2I or FCMD in humans, with early-onset muscular dystrophy as well as eye, neuromuscular junction, and CNS abnormalities (Figure S3). When treated with the biAb, the dystroglycan complex was not disrupted as observed in immunoprecipitation assays of muscle lysates from mice that had been treated with the biAb, or the parental Abs (Figure 2B).
As a proof of concept, LARGEmyd-3J homozygous mice were injected with the biAb into the tibialis anterior (TA) muscle; the left TA muscles were injected with the biAb on days 1 and 4, while the right TA muscles were injected with the parental Abs as controls. On day 5, Evans blue dye (EBD) was injected intraperitoneally into the mice, and the animals were then subjected to a forced treadmill exercise to induce muscle damage. As shown in Figures 2C and 2D, the EBD-positive fiber counts in the biAb-treated TA muscles were significantly (~50%) less than those in the control-treated TA muscles, suggesting that treatment with the biAb conferred a measure of protection against exercise-induced damage. We interpret that this was due to biAb-mediated reconnection of laminin-211 to βDG. We also confirmed that administering a mixture of the parental Abs to either laminin-211 or βDG did not exhibit a dominant-negative effect by conducting the same experiment in wild-type mice (Figure 2C).

**Systemically Delivered Abs Are Localized in the Muscle Sarcolemma**

With protein replacement therapies, biotherapeutics are typically administered to patients via intravenous infusions. Hence, for this concept to be efficacious in patients with α-dystroglycanopathy, systemic delivery of the biAb needs to result in localization of the agent in a majority of muscle tissues, particularly the sarcolemma, and the biAb needs to be sufficiently stable in situ. To evaluate this, mice were injected via the tail vein with either the biAb or a mixture of the two parental Abs (as a control), and the concentration of the Abs in the blood, muscle, and non-target tissues was then measured over 14 days. Blood samples were collected after 3 h and on days 1, 2, 3, 5, 7, and 14 post-injection for assessment. Target muscle tissues (quadriceps, TA, diaphragm, and heart) as well as non-disease-relevant tissues (lung, liver, kidney, and spleen) were also collected on days 4, 7, and 14 post-injection (following perfusion) to assess biodistribution and in situ stability. As shown in Figure 3A, both the parental Abs and the biAb were still detectable in blood on day 14, albeit at low levels, with a half-life of approximately 30 h (Table S2). The parental Abs and the biAb were also detected in various muscle groups, such as the quadriceps, TA, diaphragm, and heart for up to 14 days (see Figure 3B, showing diaphragm and quadriceps as representative examples), indicating that the tissue half-life of the biAb was longer than that in the blood. This suggested that, after the Abs reached the muscle tissues and bound their targets (laminin-211 in the basal lamina and βDG on the sarcolemma, or both in the case of the biAb), the Abs persisted for a sustained period. In contrast, Abs were barely detectable in non-target tissues. There was little to no signal in the liver or spleen on day 4, the earliest collection time (Figure 3C), suggesting that the longer half-life of the Abs in the muscle was due to binding to their respective cellular targets.

**Systemic Delivery of the biAb Targeting Laminin-211 and βDG into LARGEmyd-3J Mice Improves Muscle Function and Confers Protection from Exercise-Induced Damage**

Next, we assessed whether systemic delivery of the biAb targeting laminin-211 and βDG protected the muscles of the LARGEmyd-3J mice from exercise-induced damage and, more importantly, improved their function. As the homozygous LARGEmyd-3J mice (9–13 weeks) presented with phenotypic variability, several parameters comprising age, hindleg splay score, grip strength, wire-hang time, baseline serum creatine kinase (CK) level without treadmill exercise, and CK level after forced treadmill exercise (Figures S3A–S3F) were used to randomize the mice into treatment groups with similar parameters. After randomization, groups of LARGEmyd-3J mice were treated with the biAb or a mixture of the parental Abs or saline as controls. Wild-type mice were also administered the biAb to assess the effect of the biAb on normal muscle function.

Following biAb treatment, LARGEmyd-3J mice showed a noticeable improvement in grip strength starting at 2–3 weeks (Figure 4A), and this improvement continued until ~5 weeks post-treatment,
when it appeared to reach a plateau. biAb-treated LARGEmyd-3J mice (at the end of the study) attained nearly 70% of the grip strength noted in the wild-type mice, compared to the parental Ab- or saline-treated control mice that maintained ~55% of the grip strength noted in wild-type mice. In the wire-hang test (Figure 4B), a trend toward improvement was noted in the LARGEmyd-3J mice after 2–3 weeks of treatment with the biAb, with a significant change attained after 5 weeks. The magnitude of the improvement in the wire-hang test was not as large as that in the grip strength test. Statistical significance was attained due, in part, to the continual decline in the performance of the control LARGEmyd-3J mice. There was no significant difference in body weight between groups (Figure 5A).

The mice were subjected to the forced treadmill exercises only at the start (baseline) and the end (7 weeks of treatment) of the study to minimize muscle damage and fatigue that might have confounded the results of the other behavioral tests. As shown in Figure 4C, the untreated LARGEmyd-3J mice were only able to run for ~4 min at the start of the study, and endurance decreased further over time. In contrast, the animals treated with the biAb showed an improved run time of ~8 min at the end of the study, which represented a 2-fold increase over that noted for the control mice. We attribute these improvements in both the behavioral and muscle function tests to biAb-mediated re-engagement of laminin-211 with βDG. To assess whether treatment with the biAb also conferred some measure of protection against exercise-induced damage, we used the EBD assay. As shown in Figure 4D, the biAb-treated LARGEmyd-3J mice exhibited significantly lower numbers of EBD-positive fibers in the TA muscle, suggesting that the biAb provided some protection against exercise-induced muscle damage. However, the number of EBD-positive fiber counts in other muscle types, such as the force-bearing quads, did not show an improvement with treatment (data not shown). It is possible that, with these force-bearing muscles, excessive running on the treadmill may have induced additional muscle damage in the treated animals. We also observed no improvement in CK levels (Figure S4B), which could also be explained by this excessive running. Consistent with this premise is the finding that the histopathological changes in the quads of biAb-treated and control-treated mice were not significantly different (Figure 4E). It is also possible that the lack of improvement in the quads was due to the relatively short duration of treatment that resulted in the delivery of suboptimal concentrations of the biAb to these muscles. It should also be noted that the LARGEmyd-3J mice exhibited only modest histopathological changes between 5 and 10 months of age that hampered observations of small improvements (Figure S3J). This lack of histopathological changes could be specific to the LARGEmyd-3J model.

Unlike the physiological binding of laminin-211 to zDG O-linked matriglycans, which is multivalent,29 that of the biAb between laminin-211 and βDG is monovalent. Moreover, the binding avidity of laminin-211 to the matriglycans (~6 nM) is significantly higher than that of biAb to laminin-211, which, we determined, is at ~30 nM (Table S1).15 Thus, the overall binding of laminin-211 to dystroglycan mediated by the biAb was anticipated to be much weaker than under normal physiological conditions. In addition, it is not known whether binding of laminin-211 to the integrins, proteoglycans (heparin), or glycolipids (sulfatides) may also contribute to protecting the muscle membrane from exercise-induced damage.
These differences may limit the translatability of this strategy to patients. While a biAb-mediated therapy may improve muscle function (patients attain greater physical strength), because it does not confer protection from exercise-induced muscle damage, the benefit may be modest and short-lived.

To address some of the aforementioned limitations, we generated a biAb in a different format (e.g., CODV1g) that limits the confounding issue of steric hindrance. Unfortunately, the CODV1g construct was limited by production issues and could not be made in quantities to support in vivo testing. Thus, we also evaluated the Duobody format and were successful in generating amounts of the biAb to support testing in animals.

Efficacy of Duobody biAb Is Dose Dependent and Is Sustained for 2 Weeks Post-treatment

The selection of the 30 mg/kg dose used in the studies described earlier was based on pilot studies showing evidence of efficacy at this dose (data not shown). As the biAb (in TBTI format) used in these studies demonstrated lower affinity (due to steric hindrance) to laminin-211 than the parental Ab (Figure S2B, left panel), we elected to engineer an improved version based on the Duobody format (biAb-2) with higher affinities to both laminin-211 and βDG. This biAb-2 was used in a separate dose-response study in LARGEmyd-3j mice. We tested two doses: 5 mg/kg (which was anticipated to be ineffective based on earlier pilot studies) and 30 mg/kg. As shown in Figures 5A and 5B, the mice administered 30 mg/kg of biAb-2 demonstrated improved muscle function at 3 weeks post-treatment, while those treated with 5 mg/kg showed no change when compared to the control mice that were administered saline or the parental Ab. No significant difference in body weight was observed between the groups (Figure S4C).

Since the biAb was detected in tissue for more than 14 days (Figure 3B), the longevity of the noted efficacy was also evaluated. Mice were treated with 30 mg/kg of biAb-2 for 3 weeks, and their performance was followed for an additional 5 weeks without additional treatments. As shown in Figures 5C and 5D, the biAb-2-treated animals demonstrated slow, progressive declines in grip strength and wire-hang time following treatment cessation; however, statistically significant improvements were maintained (in both tests) for 2 weeks after treatment cessation. CK levels appeared to decrease with 30 mg/kg of biAb treatment; however, this positive trend was not statistically significant (Figure S4D). At 5 weeks after treatment cessation, the performance of the biAb-2-treated mice had returned to the baseline values noted before the start of the treatment.

It has been shown that AAV9-mediated expression of LARGE (under the transcriptional control of a muscle-specific promoter) is efficacious at correcting the aberrant O-glycosylation on zDG and ameliorating the muscle pathology in LARGE-deficient mice. We have independently confirmed these observations using a similar AAV vector. In Figure S5A, we show that glycosylation was restored above wild-type levels with AAV treatment. Using the extensor digitorum longus (EDL) muscle eccentric contraction damage assay, we further demonstrated that the EDL muscles of AAV9-LARGE1-treated mice were fully protected from eccentric contraction-induced damage, a finding that we did not observe with the biAb-2 in short-term studies. Thus, a longer term study (10 weeks of treatment) with biAb-2 was performed to determine whether accumulation of higher amounts of biAb in EDL may confer greater protection. As anticipated, an improvement in grip strength was noted that was sustained over the entire treatment period (Figure 6A); however, the magnitude of improvement was less than that seen in animals administered AAV9-LARGE1. At the end of the study, a subset of mice treated with AAV9-LARGE1, biAb-2, or the parental-Ab-treated mice were subjected to the EDL assay. As shown in Figure 6B, 4 of
the 6 EDLs from biAb-2 treated mice showed improved protection (albeit to differing extents), compared to none in the parental-Ab-treated mice. As expected, full protection of EDL muscles was observed in all the AAV9-LARGE1-treated animals. Prior to the contraction assay, we observed no change in twitch or tetanic force in the biAb-treated animals compared to the parental-Ab-treated mice (Figures S5B and S5C). This result is consistent with the results from the EBD-positive fiber assay and suggests that improving the affinity of the biAb and increasing the duration of treatment with the biAb may confer greater protection from exercise-induced muscle damage. Histopathological analysis demonstrated that significant improvement was attained with AAV9-LARGE1 treatment, but no noticeable change was observed with biAb-2 treatment (data not shown). This seems to be contradictory to the results of the EDL assay in which muscle damage protection was attained in some of the animals, albeit to a lesser degree, compared to AAV9-LARGE1. We speculate that this could be due to a combination of factors, including the time needed for biAb accumulation in muscles to a critical threshold concentration necessary to effectively and robustly protect the muscle; the time required for muscle repair to complete under normal conditions is estimated to be ~3 weeks. By contrast, the onset of LARGE1 expression from AAV9-LARGE1 occurred rapidly (within a few days) and was, therefore, able to more rapidly restore the matriglycan on zDG to confer protection from muscle damage. In addition, the ratio of biAb-mediated binding of laminin-211 to βDG was only 1:1, which is much less than that of matriglycan to laminin-211. Finally, the affinity of the biAb to its ligands was less than that with matriglycan. Thus, further improvements to the affinity and binding molar ratios (e.g., via multi-specific antibodies) could be explored to extend our findings of improved muscle function to histopathological normalization and long-term benefits in the clinic.

**DISCUSSION**

Intuitively, the genetic heterogeneity of zDG-related muscular dystrophies would require a personalized approach to managing each of the 18 causative genes identified thus far. Indeed, gene augmentation therapy to reconstitute the expression of the respective gene products has demonstrated success in several murine models of this group of diseases. However, challenges associated with developing systemic gene therapy for muscular dystrophies remain. These challenges include the relatively low efficiency of current gene delivery vectors (such as AAV vectors) in transducing the mass of affected muscles and the presence of pre-existing neutralizing Abs against the viral vectors. Both factors limit the number of eligible patients amenable to this approach. Nevertheless, innovative approaches continue to be sought to improve the tropism and efficiency of AAV vectors. While developing a separate gene delivery vector for each of the diseases is rational, this is arguably inefficient and offers a high cost-to-benefit ratio, particularly given the low prevalence of patients associated with each subset of diseases. For this reason, we rationalized that a more generalized approach that could ameliorate several subtypes of z-dystroglycanopathies might be more advantageous. Toward this aim, we engineered biAbs that could act as a surrogate molecular linker to reconnect the generalized loss of binding between laminin-211 and zDG in the LARGE<sup>myd<sup>−</sup>3</sup> mouse model. This approach could potentially treat other subtypes of z-dystroglycanopathies; however, additional animal models need to be tested.
The preclinical data generated in the LARGE<sup>myd-3F</sup> mice using the biAbs that we developed suggest that the biAbs were able to engage both laminin-211 and βDG simultaneously (see Figures 4, 5, 6, and S4A for potential mechanisms of action). Only the biAbs, not the parental Abs (administered as a mixture that engaged their targets separately), conferred a beneficial outcome (Figures 4, 5, 6, S6A, and S6B), suggesting that biAbs engaged both targets concurrently. However, attempts to visualize this physical interaction by immunofluorescence imaging were unsuccessful due to the close proximity of the sarcolemma to the basal lamina (Figure 3B). Thus, higher resolution and more sensitive physical and biochemical approaches are likely needed to address this specifically.

Evidentiary demonstration of efficacy was noted in biAb-treated LARGE<sup>myd-3F</sup> mice using validated muscle function assays. The improvements were highly significant but did not reach the levels noted in healthy animals. Moreover, with some of the tests (e.g., wire-hang tests), maximal benefit was realized after 5 weeks of treatment, and performance did not improve further with additional biAb supplementation. Possible reasons for this include the generation of anti-drug Abs; indeed, low anti-biAb Ab levels were detected after 4 weeks of treatment (data not shown). Another, and perhaps more likely, possibility is the inability of the biAb (in its current configuration) to correct the defects at neuromuscular junctions (NMJs), as this requires the restoration of the linkage between agrin and dystroglycan. The inability of the biAb to fix the abnormal NMJs noted in the LARGE<sup>myd-3F</sup> mice (Y.Z., N.G., Z.L., and D.S.B., unpublished data) may have contributed to the confounding behavioral data. Indeed, the hindleg splay assay, which requires proper functioning of the peripheral nerves, did not improve with treatment with the biAbs. Concordant with this premise, McKee et al. recently reported on the ability of a chimeric protein comprising the N-terminal domain of laminin α2 subunit fused to a nidogen fragment (that binds to the laminin γ1 subunit) to repair a defect in laminin-211 polymerization in Lama2<sup>dy-2j</sup> mice. While the forelimb strength of the treated Lama2<sup>dy-2j</sup> mice was completely restored, that of the hindlegs was only minimally improved, likely due to the lack of correction of the amyelination of the hindlimb nerves. When a grip strength test was performed with all four limbs (as in our study), the maximal improvement attained was approximately 50% that of the wild-type mice, which was similar to what we observed in the LARGE<sup>myd-3F</sup> mice. If this explanation is true, this represents an anticipated limitation of the biAb strategy, because it will not correct the binding of DG to ligands other than laminin-211, such as agrin, neurexin, and pikachurin. biAb can correct the muscular dystrophic aspects of patients with α-dystroglycanopathies but not their neurological or ocular abnormalities. In this regard, we posit that this strategy, if successful in humans, would be most applicable for α-dystroglycanopathy patients with no or moderate CNS involvement, such as those with LGMD21 and related disorders. However, it should be noted that the structural brain defects in patients with the most severe α-dystroglycanopathies (e.g., WWS or MEB) are incurred <i>in utero</i> and, thus, may not be amenable to postnatal correction by any means, even with AAV gene therapy. Nevertheless, an improvement in muscle function in the severely affected patients with CNS involvement may still offer consequential benefits to reduce caregiver burden and extend the patients’ lives.

Given the significant but incomplete correction of muscle function in this study, complementary therapies may be needed to confer a more transformative change in patients with α-dystroglycanopathies. These could include, for example, the combined use of the biAb with follistatin or anti-myostatin to enhance muscle regeneration. Hence, it may be that the biAb represents but one of a cocktail of drugs that could be used for this group of disorders. Greater improvement may also be realized if the mice are treated at an earlier age, before there is significant and perhaps irreversible damage, or with a multi-functional antibody with multi-valency binding to laminin-211 that may offer better binding and muscle damage protection. Finally, our novel strategy of using a biAb as a molecular linker to reconnect structural aberrations could be considered for related or adjacent neuromuscular diseases.

MATERIALS AND METHODS
Expression of Human and Mouse LG5 or LG4/5 Domains of Laminin α2 Subunit and βDG ECD
The LG domains of the human and murine α2 chain of laminin-211 and the ECDs of human and murine βDG were expressed in HEK cells to ensure proper folding and glycosylation. The boundaries of the LG domains were determined based on the publications of Talts et al.<sup>40,41</sup> The LG5 domains were codon optimized for mammalian expression and synthesized with an N-terminal AviTag and an HPC4 tag. Constructs were subcloned into a mammalian expression vector and used to transfect Expi293F cells using ExpiFectamine reagent (Thermo Fisher Scientific). After 7 days, the soluble protein was purified from the supernatant with an anti-HPC4 Ab that was amine coupled to an agarose resin. Alternatively, LG4/5 domains were codon optimized and paired with an mlgG2a Fc fusion partner and a TEV cleavage site to allow for removal of the IgG domain. After synthesis, each construct was subcloned into a mammalian expression vector and was used to transfect Expi293F cells using ExpiFectamine reagent (Thermo Fisher Scientific). After 7 days, soluble protein was purified from the supernatant with a HiTrap MabSelect SuRe column (GE Healthcare). When necessary, the fusion partner was cleaved off using TurboTEV enzyme (Nacalai USA, San Diego, CA, USA) and removed by purification using Ni-NTA resin (QIAGEN).

To express the ECD of murine βDG, an <i>Escherichia coli</i> codon-optimized cassette encoding the murine βDG ECD and a C-terminal HPC4 tag was synthesized and subcloned in frame to an N-terminal maltose binding protein (MBP) fusion partner in a Novagen pET22b vector. The vector was used to transform Origami(B(DE3)pLysS cells (Novagen). Protein expression in bacteria was induced with 1 mM isopropyl β-D-1-thiogalactopyranoside. For protein purification, cell pellets were resuspended in 50 mM Tris-HCl, 150 mM NaCl (pH 7.4) containing EDTA-free protease inhibitors (Roche) and lysed by sonication. The lysate was clarified by centrifugation and filtration.
and then processed on an amylose resin (New England Biolabs) equilibrated with 50 mM Tris-HCl, 150 mM NaCl (pH 7.4). The fusion protein was eluted using the same buffer containing 10 mM maltose. Calcium (final concentration, 1 mM) was added to the amylose elution, and the fusion protein was purified further on an anti-HPC4 affinity column. Alternatively, βDG ECD was codon optimized and paired with an mLgG2a Fc fusion partner and a TEV cleavage site to allow for the removal of the IgG domain as described for LG4/5 domains.

**Ab Generation**

**Mouse Immunization**

BALB/c mice were immunized with human βDG, human LG-5, or human LG-4/5 and then boosted with these proteins 3–4 times every 2 weeks. Mice immunized with human LG-4/5 were additionally boosted three times with human merosin every 2 weeks and once with a synthetic peptide with a sequence identical between human and mouse LG-5 (amino acid sequence GPRAVGGFKV GLDLLVEFE).

**Hybridoma Generation**

Hybridoma cells were made by fusing mouse myeloma cells (from the BALB/c B-lymphoblast cell line SP2/0 fused with Sendai virus) deficient in adenosine phosphoribosyltransferase (APRT) with spleen cells from immunized mice. Hypoxanthine, azaeagine, and thymidine (HAT) selection and serial dilutions were performed to achieve single-cell clonality.

**Design of biAb Recognizing LG4/5 and βDG ECD**

A biAb recognizing both LG5 and βDG was designed with one set of anti-LG5 V\(_{\text{L}}\) and V\(_{\text{H}}\) sequences and one set of anti-βDG V\(_{\text{L}}\) and V\(_{\text{H}}\) sequences in two different anti-LG5 and βDG orientations, resulting in two biAb constructs (anti-LG5 × βDG and anti-βDG × LG5) in the TBTI format (Figure 2A). Briefly, the V\(_{\text{H}}\) sequences of a pair of binders were fused with a linker (GGGGSG), and then fused either to human IgG1 (huIGHG1; GenBank: Q569F4) or murine IgG1 (muIGHG1; GenBank: AAA75163.1) heavy-chain Fc. The V\(_{\text{L}}\) sequences of paired binders were fused to the constant human kappa chain (huIGKC; GenBank: BAB33404). An improved biAb (biAb-2) was also created with higher affinity Abs to laminin-211 and βDG in an alternative biAb Duobody format\(^{29}\) that was not subject to steric hindrance of the inner V\(_{\text{L}}\) and V\(_{\text{H}}\).

**Construction of Expression Plasmids for the Production and Purification of Recombinant mAbs and biAbs**

The light-chain and heavy-chain DNA fragments of the parental mAbs and biAbs were cloned into the episomal expression vector pXL, an analog of the pTT vector described by Durocher et al.,\(^{43}\) creating plasmids for transient mammalian expression of biAbs in Expi293 cells. The mAbs and biAbs from cell culture media were purified using a HiTrap MabSelect SuRe column (GE Healthcare) following the manufacturer’s protocol, eluted with 0.1 M citrate (pH 3.0), and neutralized immediately with 1 M Tris-HCl (pH 8.0). The mAbs and biAbs were buffer exchanged into PBS, filter sterilized (Milli-Q 0.22-µm syringe filter, SLGV033RS), and used for in vitro characterization and in vivo applications.

**Dot Blot, SPR, and ELISA Characterizations of mAbs and biAbs**

**Dot Blot.** Various antigens (BioLamina) used for dot blotting were serially diluted in PBS at 5× per dilution, typically with the highest amount at 50 ng total protein and the lowest at 0.08 ng. Five micro-liters of each sample was placed onto a nitrocellulose membrane (Bio-Rad) with heavy filter paper using a 20-µL multichannel pipette. After the membranes were air dried for 15 min, the membranes were processed as in a regular western blot. Membranes were blocked by incubation with 5% (w/v) non-fat milk (Bio-Rad) in Tris-buffered saline (TBS) with 0.05% Tween-20 (TBS-T) for 1 h at room temperature, and then primary Abs (typically at 1 µg/ml) in 5% low-fat milk were added to the membrane and mixed gently for an additional 1 h at room temperature. Membranes were washed three times in TBS-T for 5 min each. Goat anti-mouse IgG-HRP (horseradish peroxidase) (Pierce, 1:5,000 dilution) or goat anti-human IgG-HRP (Millipore, 1:5,000 dilution) in 5% milk was added and incubated with membranes for another 1 h, followed by three washes in TBS-T as described earlier. Blots were developed with SuperSignal Chemiluminescent Substrate (Pierce) for 5 min and exposed to film for various amounts of time ranging from 1 s to 10 min.

**ELISA.** A sandwich ELISA was developed by taking advantage of the difference in Abs binding between mouse and human IgG Fc (Figure S2C). First, 96-well plates were coated with 100 µL (0.5 µg/mL) LG5 in PBS overnight at 4°C, after which the coating solutions were removed. The plates were then blocked with 5% fetal bovine serum in PBS for 1 h, followed by three washes with PBS. A 100-µL aliquot of biAb (or a single parental Ab to laminin-211 or βDG) with the human IgG backbone (anti-laminin α2 × βDG) at 10 µg/mL was added to each well and incubated at room temperature for 2 h with gentle shaking. Thereafter, the plates were washed three times with PBS, and various concentrations of βDG ECD fused to mouse Fc (βDG ECD-mFc) in 100-µL aliquots were added to the wells in 2-fold serial dilutions, followed by incubation for 2 h with gentle shaking at room temperature. After three washes with PBS, HRP-conjugated anti-murine Fc secondary Ab was added and incubated for 45 min. The plates were washed four times with PBS and developed for colorimetric measurements. Signals could be detected only when both LG5 and βDG ECD-mFc were added in the assay. No signal was detected when βDG ECD-mFc was omitted or when only one parental Ab was added. This procedure could be reversed using βDG for coating followed by LG5-mFc with similar results (data not shown).

**SPR Analysis of Abs.** A Biacore T100 kinetics/affinity assay was optimized to capture the Abs (mAbs and biAbs) on chips and screen for ligand-binding affinities. Briefly, a Biacore human or mouse Ab capture kit (GE Healthcare) was used to immobilize ~10,000 RU of mouse anti-hIgG or rabbit anti-mIgG on a CMS Series S biosensor using amine-coupling techniques. Abs and a negative isotype control
were diluted to 20 μg/mL in HBS-EP+ running buffer (10 mM HEPES, 150 mM NaCl, 3 mM EDTA, 0.005% polysorbate 20). The Abs were captured for 60 s at 10 μL/min, and human and murine LG5 or LG4/5 or human βDG and MBP-βDG ECD fusion proteins were diluted to 80, 40, 20, 10, 5, 2.5, 1.25, and 0 nM in HBS-EP+ running buffer. Analytes were injected for 5 min to measure association, followed by 10 min of buffer for dissociation at 30 μL/min. The Ab capture surface was regenerated between cycles using a 60-s injection of 12 mM HCl at 30 μL/min. Reference-subtracted sensograms were fitted to a 1:1 binding model using Biacore T100 Evaluation software v.2.0.2. The process of sequential binding of LG5 and βDG to Ads on Biacore was essentially the same as described earlier, except that a 100-nM solution of one analyte was injected for 2 min at 30 μL/min, immediately followed by injection of the second analyte, also at 100 nM, to detect the second binding event (Figure S2B).

Animals
The LARGEmyd-3J/GrsrJ mice (stock #008581) were purchased from the Jackson Laboratory after custom cryo-recovery and breeding between heterozygous males and females. A genotyping method was developed for the myd-3j mutation (Y.Z., N.G., and Z.L., unpublished data). Homozygous mice were confirmed by phenotypic screenings, and both male and female mice were used for the studies in this paper. Homozygous mice generally began to exhibit symptoms as early as weaning. They initially displayed an inability to splay their hindlegs outward when held up by the tail, progressing to a swaying gait and were eventually dragged of their hindlegs. Homozygous LARGEmyd-3J/GrsrJ mice (stock #008581) were purchased from the Jackson Laboratory after custom cryo-recovery and breeding between heterozygous males and females. A genotyping method was developed for the myd-3j mutation (Y.Z., N.G., and Z.L., unpublished data). Homozygous mice were confirmed by phenotypic screenings, and both male and female mice were used for the studies in this paper. Homozygous mice generally began to exhibit symptoms as early as weaning. They initially displayed an inability to splay their hindlegs outward when held up by the tail, progressing to a swaying gait and were eventually dragged of their hindlegs.

Biodistribution of mAbs and biAb in Mice
Wild-type mice at ~8 weeks of age were injected intravenously (i.v.) via the tail vein with 30 mg/kg of the biAb or a mixture of parental Abs (15 mg/kg each) with the mlgG1 backbone. Blood serum samples were collected by submandibular vein bleeding at 3 h; 24 h; and 2, 3, 4, 7, and 14 days after administration. The parental Ab or biAb level at each time point was measured using an ELISA developed in house. Briefly, a 96-well plate was coated with 100 μL per well of mouse MBP-βDG at 0.5 μg/mL overnight at 4°C. The plate was washed with PBS and blocked with blocking buffer containing 5% BSA for 1 h followed by 3 washes with PBS; and 100 μL of each serum sample at various dilutions was then added into the wells. The plates were incubated for 2 h with gentle shaking at room temperature and washed 4 times with PBS. HRP-conjugated goat anti-mouse IgG was added and incubated for 45 min. After 6 washes with PBS, the plate was developed and measured via colorimetry. Purified biAb solutions in assay diluent at various concentrations were used as standards. Blood half-life of antibodies was estimated according to the graph in Figure 3A.

To determine tissue biodistribution, four mice per group were perfused and sacrificed at days 4, 7, and 14. Various muscle tissues and non-target tissues were harvested in optimum cutting temperature (OCT) compound (EM Bioscience), snap-frozen in isobutane/dry ice, and stored at −80°C until use. To stain for the mAbs and biAb in tissues, tissues were cryo-sectioned at 10-μm thickness, air-dried unfixed, and stained with fluorescently labeled anti-mlgG Fc Ab.

Local Injection of biAb into the TA Muscle of the LARGEmyd-3J Mice and Analysis of Muscle Damage
A group of LARGEmyd-3J mice (n = 10) were given intramuscular injections into the left and right TA muscles. The left TA muscle received two injections of biAb-mlgG1 (at 0.7 μg/μL in 50 μL saline per injection with 29G needles) 3 days apart. The right TA muscle received two injections of 50 μL parental Ab mixture per injection. One day after the last intramuscular injection, all the mice received intraperitoneal injections of EBD (at 10 mg/mL, with 50 μL given per 10 g body weight). Eighteen hours after intraperitoneal injection of EBD, all the mice were forced to exercise on a treadmill (Columbus Instruments) until exhaustion. After euthanasia and perfusion, the TA muscles were removed, cut, and placed in OCT. The tissue was cryo-sectioned at a thickness of 10 μm at four different levels in triplicate, 100 μm apart. For staining, the slide sections were quickly dipped into cold PBS and fixed in ice-cold acetone for 15 min. Slides were washed (3 × 5 min) in PBS, outlined with a hydrophobic marker, and blocked with 2% BSA with 1:100-diluted normal goat serum in PBS overnight at 4°C. The next day, the slides were incubated with anti-mouse IgG Alexa Fluor 488 (Invitrogen) at 1:400 dilution for 2 h at room temperature. Slides were washed 3 times (5 min each) with TBS-T and then washed 1 time for 5 min with PBS and mounted with VectaShield mounting medium with DAPI. Slides were sealed with coverslips and visualized with an inverted microscope (Olympus IX71) using appropriate filter sets. Pictures were acquired using a charge-coupled device (CCD) Hamamatsu digital camera (C8484) and MetaMorph software (v.7.8.1.0). Each field was exposed for 1 s for Alexa Fluor 488 and EBD or 0.5 s for DAPI. The pictures were subsequently overlaid using Adobe Photoshop CS5.

For EBD-positive fiber counting, tissue sections were processed as described earlier. The EBD-positive fibers in each section were counted manually twice at each of the four levels for both the left and right TA muscles of each animal. The numbers of EBD-positive fibers in the sections at four levels were averaged to yield the EBD-positive fibers per TA muscle sample. In every animal analyzed, there were fewer EBD-positive fibers in the biAb-injected left limb than in the parental Ab-injected right limb. The difference between the left and right TA muscles reached statistical significance (p < 0.05) as determined by a paired Student’s t test.

Method for IP
For each sample, 90 mg frozen muscle tissue that was locally injected with saline, parental Ab, or bispecific antibody as described earlier was placed into a 2-mL bead beater tube together with 1.8 mL ice-cold immunoprecipitation (IP) lysis buffer (TBS [pH 8], 1% NP40,
0.5% sodium deoxycholate, 10 mM EDTA, 10 mM EGTA, PMSF, and protease inhibitors) and one sterile 5-mm steel bead. The samples were subjected to 2 cycles (10 min per cycle) in the bead beater (Tissuelyser II, QIAGEN) at maximum speed (30Hz) at 4°C, after which the samples were removed and then slowly rotated for an additional hour at 4°C. Samples were then centrifuged at 13,000 rpm for 10 min at 4°C, and the supernatants were removed and aliquoted (~900 μL). For direct IP of DGC, an anti-βDG antibody (Leica clone 43DAG1/8D5), which has a different epitope than that in our biAb treatment, was utilized. For in vivo bound IP, no additional antibody was added. The supernatant generated as described earlier was incubated overnight via slow rotation at 4°C. Protein A/G bead slurry was then added and further incubated overnight via slow rotation at 4°C. The resulting mixture was centrifuged (2,000 rpm for 5 min), and the supernatant was removed. The resulting pellet was washed extensively with TBS, re-suspended in 120 μL SDS-PAGE loading buffer with fresh β-mercaptoethanol, boiled for 10 min, and centrifuged (5,000 rpm for 5 min). The supernatant was assessed by western blot using the polyclonal AF6868 DAG1 antibody (R&D Systems).

**Systemic Delivery of biAb into LARGE™**

**Mice and Analysis of Muscle Function Improvement and Damage Protection**

To ensure that the LARGE™ mice (between 9 and 13 weeks of age) started with a similar performance baseline prior to testing, several parameters were measured, including hindleg splay, grip strength, wire-hang time, and treadmill run time as well as CK levels (Figure 4). Based on these test results, the LARGE™ mice were randomized into three groups (n = 16) with similar average scores. One group received the biAb (30 mg/kg), and the other two groups received the parental Abs (a mixture of anti-laminin-211 at 15 mg/kg and anti-βDG at 15 mg/kg) or saline as a control. As an additional control, another group of wild-type mice (n = 6) received the same dose of biAb. The mice were dosed i.v. via tail vein injection twice a week for a total of 7 weeks. One day after the last injection, all the mice were tested for wire-hang time, grip strength, and hindleg splay, after which they received EBD as described earlier. After waiting overnight, all the mice were exercised via a forced treadmill run until exhaustion (with 15 min as the cutoff). One hour after treadmill running, the animals were euthanized with CO₂ and perfused with PBS, and various muscles were collected and placed in OCT compound. The muscles were cryo-sectioned and processed for EBD-positive fiber counting as described earlier. Statistical analysis was performed with one-way ANOVA followed by Tukey’s multiple comparison test. A similar study with older mice (between 19 and 22 weeks of age) was repeated for confirmation, with two groups of LARGE™ mice receiving the biAb or saline and a wild-type group receiving saline. For the treatment withdrawal study, mice were dosed with 5 or 30 mg/kg of the biAb, parental Abs, or saline for 3 weeks, when a significant improvement on grip strength and wire-hang time was observed. Then, treatment was stopped, but behavioral tests were continued weekly for the 30 mg/kg biAb group, the saline control group, and the wild-type group until they reached baseline or control levels. The 5 mg/kg and parental Ab groups were sacrificed for tissue collection immediately after 3 weeks of treatment.

**Behavioral Tests**

For the hindleg splay test, the mice were lifted by their tails, and the positions of the hindlegs relative to the body were recorded and graded. The wire-hang test was conducted by placing mice on a wire grid and letting them acclimate for 1 min; then the wire grid with the mouse grasping on was slowly turned upside down in 2 s, and the time the mouse held onto the grid was recorded, with a cutoff time of 60 s. The test was repeated 3 times for each mouse, and the results were averaged. Grip strength was evaluated by placing the mice on a grip strength meter (Columbus Instruments), allowing the mouse to grasp the metal grid firmly and pulling the tail horizontally until the mouse let go; then, the force was recorded. The test was repeated 5 times, with a 1-min rest between trials. The highest and lowest readings for each mouse were removed, and the results presented are the average of the three remaining readings. For treadmill run time, mice were placed onto individual lanes of a treadmill equipped with an electric shocking grid (model 1055-SRM Exer-3/6, Columbus Instruments). The animals were acclimated to the treadmill for 5 min, and then the mice were tested with a defined protocol with increasing speed. When a mouse spent more than 3 s on the shocking grid without being able to run, the shocking grid was turned off, and the total run time was recorded. All behavioral tests were performed by researchers blinded to mouse identity and treatment, with the results revealed after testing.

**Ex Vivo EDL Force Measurement**

Contractile properties of EDL muscles isolated from wild-type and LARGE™ mice were measured by using a 300B dual-mode servomotor transducer and DMC/DMA software (Aurora Scientific), as described elsewhere. Briefly, the distal tendon of the EDL was tied to the servomotor by using a 6-0 silk suture, and the proximal tendon was clamped to a fixed post. The EDL muscles were immediately immersed in an oxygenated bath (95% O₂, 5% CO₂) that contained modified Krebs-Ringer buffer at 25°C. Muscle length and the current of stimulation were adjusted until an isometric twitch contraction achieved the maximum force (induced by a single stimulus with a pulse duration of 0.2 ms). After the optimization, maximum twitch forces were obtained by a single stimulus with a pulse width of 0.5 ms. At the optimized muscle length (Lo), maximum isometric tetanic forces were determined by applying tetanic stimulation for 500 ms at a frequency of 120 Hz with a pulse width of 0.5 ms while the muscle was lengthened by 7.5% Lo at 0.5Lo/s during the last 200 ms. The muscles were rested for 2 min between successive contractions. Maximum tetanic force produced before the lengthening contraction was obtained at each cycle to calculate the force drop as a percentage of the force obtained during the first cycle.

**Data Availability**

The datasets generated during and/or analyzed during the present study are not publicly available per Sanofi intellectual property policy
Y.Z. conceived the therapeutic strategy, designed the experiments, and interpreted and analyzed the data. N.G., W.H.B., and S.H.C. supervised the work. Y.Z. wrote the manuscript, and N.G., W.H.B., S.H.C., and L.M.S. edited the manuscript.

CONFLICTS OF INTEREST
All the authors of this paper are present or past employees of Sanofi.

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