Expression of the dystrophin isoform Dp116 preserves functional muscle mass and extends lifespan without preventing dystrophy in severely dystrophic mice

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Dp116 is a non-muscle isoform of dystrophin that assembles the dystrophin–glycoprotein complex (DGC), but lacks actin-binding domains. To examine the functional role of the DGC, we expressed the Dp116 transgene in mice lacking both dystrophin and utrophin (mdx:utrn−/−). Unexpectedly, expression of Dp116 prevented the most severe aspects of the mdx:utrn−/− phenotype. Dp116:mdx:utrn−/− transgenic mice had dramatic improvements in growth, mobility and lifespan compared with controls. This was associated with increased muscle mass and force generating capacity of limb muscles, although myofiber size and specific force were unchanged. Conversely, Dp116 had no effect on dystrophic injury as determined by muscle histopathology and serum creatine kinase levels. Dp116 also failed to restore normal fiber-type distribution or the post-synaptic architecture of the neuromuscular junction. These data demonstrate that the DGC is critical for growth and maintenance of muscle mass, a function that is independent of the ability to prevent dystrophic pathophysiology. Likewise, this is the first demonstration in skeletal muscle of a positive functional role for a dystrophin protein that lacks actin-binding domains. We conclude that both mechanical and non-mechanical functions of dystrophin are important for its role in skeletal muscle.

INTRODUCTION

Duchenne muscular dystrophy (DMD) is a severe muscle-wasting disorder caused by mutations in the X-chromosome-linked dystrophin gene. The disease is relatively common, affecting 1 in 3500 males, and has no effective treatment (1). A wide variety of studies of disease pathogenesis have been performed using the mdx mouse model, a naturally occurring dystrophin-deficient strain (2). Muscles in mdx mice show many of the histological characteristics of early DMD muscles, including necrosis, inflammation and regenerating myofibers (3,4). Despite being an exact genetic model of DMD, the mdx mouse has been criticized for a failure to recapitulate the severe progressive nature of muscular dystrophy in humans (5). Unlike humans, mdx mice do not have a dramatically shortened lifespan, and display muscle hypertrophy and retention of muscle function throughout the majority of their life (6–8). In contrast, mice deficient in both dystrophin and its autosomal homologue utrophin (mdx:utrn−/−) display many of the severe features of DMD, including growth retardation, skeletal deformities and premature death (9,10). This observation supports the hypothesis that the upregulation of utrophin observed in mdx muscles compensates for dystrophin deficiency, leading to the mild phenotype of these mice. Overexpression of utrophin prevents muscular dystrophy in transgenic mdx mice, indicating that utrophin and dystrophin can be functionally redundant (11).

Dystrophin is required for complete assembly of the dystrophin–glycoprotein complex (DGC), which consists of the dystroglycan, sarcoglycan–sarcospan and dystrobrevin–syntrophin sub-complexes (reviewed in 12). Dystrophin interacts with costameric F-actin via its N-terminal and central

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actin-binding domains and with transmembrane dystroglycan via its WW and cysteine-rich domains, creating a mechanically strong link between the extracellular matrix and the cytoskeleton (13,14). This link functions to dissipate the forces of contraction from within muscle fibers to the extracellular matrix, protecting the sarcolemma from stresses during exercise, and is referred to as the mechanical function of dystrophin (15,16).

Dystrophin also potentially serves as a scaffold for signaling complexes, as many of the proteins in the DGC bind to or are substrates for known signaling molecules, including neuronal nitric oxide synthase (nNOS), caveolin-3, Grb2 and various kinases (17–22). However, restoration of the DGC without restoring the mechanical function of dystrophin does not improve the phenotype of mdx mice (23–25). These results support the hypothesis that the loss of the mechanical function of dystrophin is the primary cause of muscular dystrophy in mdx mice. The increased severity of mdx:utrn<sup>−/−</sup> phenotype compared with mdx has been largely unexplained, but could be due to further weakening of the mechanical connection between the cytoskeleton and the extracellular matrix, caused by the additional loss of utrophin. Another possible explanation is that utrophin upregulation in mdx muscles is adequate for the maintenance of signaling through the DGC, thus the severe consequences of the double mutant are due to the loss of both the mechanical and signaling functions in muscle.

In a previous study, we described transgenic mice that express the dystrophin isoform Dp116 in skeletal muscle (25). Dp116 is normally expressed in Schwann cells of the peripheral nervous system where it associates with a DGC-like complex that, as in skeletal muscle, binds to laminin-2 in the extracellular matrix (26). Furthermore, Dp116 contains the complete dystroglycan-binding domain of dystrophin, but does not have any of the actin-binding motifs, allowing us to dissect the mechanical and signaling properties of dystrophin. Expression of Dp116 and restoration of the DGC did not improve the phenotype of mdx<sup>cav</sup> mice, consistent with previous studies demonstrating that actin-binding domains of dystrophin are required to prevent dystrophic injury. However, we hypothesized that expression of Dp116 in mdx:utrn<sup>−/−</sup> mice would allow a clearer dissection of the signaling and mechanical roles of dystrophin, as residual utrophin in mdx muscle may maintain critical signaling processes. Here we showed that the increased severity of the mdx:utrn<sup>−/−</sup> phenotype compared with that of mdx mice is not simply due to a further loss of the structural link between the actin cytoskeleton and the extracellular matrix. Instead, our data suggest that the dystrophin/utrophin glycoprotein complex itself plays an important role in maintaining muscle mass, even in the absence of a mechanically strong link to the cytoskeleton.

RESULTS

Generation of transgenic Dp116 mice on the mdx:utrn<sup>−/−</sup> background

Dp116/mdx transgenic mice were crossed with mdx:utrn<sup>−/+</sup> mice (9) for two generations to obtain Dp116:mdx:utrn<sup>−/−</sup> mice (see Materials and Methods). From nearly 300 pups in line 2197, we obtained 10.9% of the genotype mdx:utrn<sup>−/−</sup> and 7.4% of the genotype Dp116:mdx:utrn<sup>−/−</sup>, close to the expected ratio of 12.5% for each genotype. To verify that the genotypes accurately reflected expression of the three gene products of interest, we performed immunofluorescence staining of skeletal muscles using polyclonal antibodies against the dystrophin N-terminus, utrophin-A and the Flag epitope (Fig. 1). Dp116 was appropriately expressed and localized to the sarcolemma in esophagus, diaphragm, intercostal and various hindlimb muscles, although expression in the diaphragm was consistently much lower than in the limb muscles (data not shown). As expected, no significant expression was seen in cardiac muscle because we used the human α-skeletal actin promoter that selectively expresses the transgene in skeletal muscle. We have previously shown that expression of Dp116 at the sarcolemma is sufficient for restoration of the DGC, excluding neuronal nNOS (25).

Dp116 expression increases muscle mass and lifespan of mdx:utrn<sup>−/−</sup> mice

As expected, mdx:utrn<sup>−/−</sup> mice were smaller than their littermates at weaning and showed significant growth retardation throughout their lifespan. Although Dp116:mdx:utrn<sup>−/−</sup> mice were slightly smaller on average than control Dp116 transgenic mice with a functional utrophin gene (Dp116:mdx:utrn<sup>+/−</sup>), they continued to gain body mass at a similar rate until beyond 20 weeks of age (Fig. 2A). Control mdx:utrn<sup>−/−</sup> mice developed severe kyphosis of the spine, joint contractures and reduced mobility between 8 and 10 weeks of age. In contrast, Dp116:mdx:utrn<sup>−/−</sup> mice did not develop any of these overt signs of muscle weakness and were essentially indistinguishable from their littermates that expressed utrophin (Fig. 2C and Supplementary Material, Movie). Consistent with their improved physical appearance, Dp116:mdx:utrn<sup>−/−</sup> mice had a longer lifespan than the non-transgenic controls (Fig. 2B). The median lifespan of mdx:utrn<sup>−/−</sup> mice in this study was 12 weeks (n = 21, 7 males, 14 females) with the longest lived mouse reaching 20 weeks of age. In contrast, the median lifespan of Dp116:mdx:utrn<sup>−/−</sup> transgenic animals from line 2197 (n = 15, 10 males, 5 females) was found to be 46 weeks. Interestingly, none of the female mice lived beyond 46 weeks of age, while 7 of the 10 male mice lived beyond 46 weeks of age. To confirm that the effect was specific to the Dp116 transgene, we crossed the independent line 2354 onto the mdx:utrn<sup>−/−</sup> background. Thus far several mice from this line (including both males and females) have surpassed 35 weeks of age and have shown similar growth, weight gain and mobility compared with line 2197. We also observed Dp116:mdx:utrn<sup>+/−</sup> mice as controls (n = 7, all males). One of these mice developed a tumor and had to be euthanized at 42 weeks age, but the remaining 6 have remained healthy for greater than 1 year.

We sacrificed C57Bl/6, mdx, mdx:utrn<sup>−/−</sup> and Dp116:mdx:utrn<sup>−/−</sup> mice at 8 weeks of age to examine their skeletal muscles. The mass of both the tibialis anterior (TA) and quadriceps muscles was increased in Dp116:mdx:utrn<sup>−/−</sup> mice relative to mdx:utrn<sup>−/−</sup> controls, although not to the extent of corresponding mdx muscles (Table 1). The total
cross-sectional area of hematoxylin and eosin-stained sections of Dp116:mdx:utrn^{2/2} muscles was increased compared with the mdx:utrn^{2/2} controls but not compared with mdx, consistent with the increase in total muscle mass (Fig. 3A). Serum creatine kinase levels were also measured for wild-type, mdx:utrn^{2/2} and Dp116:mdx:utrn^{2/2} mice. Both mdx:utrn^{2/2} and Dp116:mdx:utrn^{2/2} mice had serum creatine kinase levels ∼10-fold higher than wild-type controls but did not differ significantly from each other (Table 1).

**Dp116 expression does not improve histopathology or mechanical properties of mdx:utrn^{−/−} muscles**

Examination of hematoxylin and eosin-stained cross-sections showed no gross improvement in dystrophic pathology in quadriceps muscles from Dp116:mdx:utrn^{−/−} mice compared with mdx or mdx:utrn^{−/−} muscles (Fig. 3). Similar pathology was found in the TA, soleus, diaphragm and intercostal muscles (data not shown). These results indicate that although muscle mass and function is preserved in Dp116:mdx:utrn^{2/2} mice, this cannot be explained by protection from dystrophic damage. Myofiber cross-sectional area was reduced to a similar extent (median cross-sectional area 37% of wild-type) in both Dp116:mdx:utrn^{−/−} and mdx:utrn^{−/−} mice and the percentage of centrally nucleated fibers remained elevated at >70% (Fig. 4A and C). The density of myofibers was measured independently to determine whether the increase in total muscle mass seen in Dp116:mdx:utrn^{−/−} was due to accumulation of non-muscle tissue or an increase in absolute number of muscle fibers. Fiber density was similar in both Dp116:mdx:utrn^{−/−} mice and mdx:utrn^{−/−} (Fig. 4B), indicating that fibrosis and the accumulation of other non-muscle tissues are not responsible for the observed increase in gross muscle mass.

Muscles from mdx and mdx:utrn^{−/−} mice display increased susceptibility to contraction-induced injury and a reduction in specific force (27,28). We evaluated mechanical properties of the TA in Dp116:mdx:utrn^{−/−} mice and compared this to both mdx and mdx:utrn^{−/−} to determine whether there was any improvement in the transgenic animals. Maximal force generation at optimal fiber length was increased by ∼50% in Dp116:mdx:utrn^{−/−} animals when compared with mdx:utrn^{−/−} mice, confirming that the increased gross muscle mass in the Dp116:mdx:utrn^{−/−} mice is largely composed of functional muscle tissue (Fig. 4D). However, specific force (force/cross-sectional area) was not significantly improved (Fig. 4E). Interestingly, Dp116:mdx:utrn^{−/−} mice demonstrated an improved resistance to contraction-induced injury that was similar to that seen in mdx mice, but remained reduced when compared with C57Bl/6 controls (Fig. 4F).

An established characteristic of mdx:utrn^{−/−} muscles is a preponderance of oxidative muscle fibers and upregulation
of slow muscle genes, which has been proposed to reflect a signaling defect that could contribute to the severe phenotype of these mice (29,30). We performed nicotinamide adenine dinucleotide (NADH) staining on sections of quadriceps and TA muscles from wild-type, mdx, mdx:utrn\(^{-/-}\) and Dp116:mdx:utrn\(^{-/-}\) mice to determine the oxidative potential of myofibers. Wild-type TA and quadriceps muscles showed a mosaic pattern of darkly stained oxidative and lightly stained glycolytic fibers (Fig. 5). Conversely, both Dp116:mdx:utrn\(^{-/-}\) and mdx:utrn\(^{-/-}\) muscles had more dark and intermediate staining showing an increased population of oxidative fibers (Fig. 5). These data show that the abnormal fiber-type distribution seen in mdx:utrn\(^{-/-}\) muscles is correlated with the severe dystrophy in these mice, but is not sufficient to cause the failure to thrive and dysfunction associated with the phenotype.

**Figure 2.** Expression of Dp116 increases viability of mdx:utrn\(^{-/-}\) mice. (A) Both male and female transgenic Dp116:mdx:utrn\(^{-/-}\) mice had continued weight gain after 10 weeks of age, while the control mdx:utrn\(^{-/-}\) mice displayed a failure to thrive beyond 10 weeks of age. However, the transgenic Dp116:mdx:utrn\(^{-/-}\) mice were consistently smaller than transgenic mice with one functional utrophin allele. Error bars signify ± SEM and are not shown for time points where only one surviving mouse remained in the cohort. (B) Expression of Dp116 increased lifespan of mdx:utrn\(^{-/-}\) mice. Kaplan–Meier analysis demonstrated a significant increase in survival \((P < 0.0001)\) of Dp116:mdx:utrn\(^{-/-}\) mice \(n = 15\), 5 females, 10 males) compared with control mdx:utrn\(^{-/-}\) mice \(n = 21\) (14 females, 7 males). Each tick-mark in the transgenic cohort indicates the age of a surviving mouse at the end of this study. (C) An 11-week-old mdx:utrn\(^{-/-}\) male is shown to demonstrate joint contractures of the hind limbs (upper left). The Dp116:mdx:utrn\(^{-/-}\) male littermate does not suffer from joint contractures and is able to extend its hind limbs (upper right). The same littermates are shown in the lower panel to demonstrate that spinal kyphosis and reduced musculature in the mdx:utrn\(^{-/-}\) control (left) are absent in the Dp116 transgenic littermate (right).

Table 1. Comparison of muscle mass and serum creatine kinase levels in age-matched transgenic and control mice

<table>
<thead>
<tr>
<th></th>
<th>C57Bl/6</th>
<th>mdx</th>
<th>mdx:utrn(^{-/-})</th>
<th>Dp116:mdx:utrn(^{-/-})</th>
</tr>
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<tbody>
<tr>
<td>TA mass (mg)</td>
<td>36.7 ± 2.0</td>
<td>65.9 ± 1.6*</td>
<td>31.9 ± 2.6</td>
<td>41.1 ± 0.90***</td>
</tr>
<tr>
<td>Quad mass (mg)</td>
<td>165.2 ± 1.5*</td>
<td>233.7 ± 3.2*</td>
<td>132.0 ± 7.4</td>
<td>158.5 ± 4.5**</td>
</tr>
<tr>
<td>Serum CK (U/L)</td>
<td>670 ± 149***</td>
<td>ND</td>
<td>8473 ± 2780</td>
<td>6575 ± 1350</td>
</tr>
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Significantly different from mdx:utrn\(^{-/-}\): *\(P < 0.001\), **\(P < 0.01\), ***\(P < 0.05\).

**Esophageal pathology may contribute to mortality in Dp116:mdx:utrn\(^{-/-}\) mice**

Despite a dramatic increase in longevity and muscle mass in Dp116:mdx:utrn\(^{-/-}\) mice, average lifespan remained reduced compared with control animals (Fig. 2B). Mice remained active and healthy for several months, but their condition began to deteriorate \(< 2\) weeks before death. The exact cause of death in these mice is unclear; however, we have repeatedly observed a grossly dilated esophagus in aged mice examined post-mortem (Supplementary Material, Fig. S1). This esophageal pathology correlated with lethargic behavior, a reduction in total body mass, and disheveled appearance prior to death (Supplementary Material, Fig. S1C). The end-stage esophagus was severely dilated compared with esophagi isolated from
8-week-old animals, indicating that gross esophageal deformities developed with age and were not present from birth in Dp116:mdx:utrn \(2/2\) mice. Dilation was most pronounced in the region of the esophagus located immediately adjacent to the diaphragmatic esophageal hiatus, above the rostral surface of the diaphragm. In addition, esophagi in aged animals were found to be impacted with solid food, bedding debris and hair (Supplementary Material, Fig. S1A). A similar pathology was not observed in terminal mdx:utrn \(2/2\) mice, nor was any similar pathology observed in Dp116:mdx:utrn \(2/2\) muscles at 8 or 12 weeks of age (Fig. 6A, Supplementary Material, Fig. S1A and data not shown).

Closer inspection of esophageal cross-sections revealed morphological changes in the esophageal walls (Fig. 6A). Both the mucosal epithelium and the muscularis externa were remarkably reduced in overall thickness. In addition, qualitative changes were apparent within the individual layers of the mucosal epithelium. The cornified epithelium was hyperkeratinized, which may represent a reactive response to abrasion, secondary to impaired transport of solid foods through the esophagus. In contrast, intermediate epithelial zones were significantly reduced or absent. The causative factors leading to esophageal impaction and dilation remain to be deciphered, but may be related to impaired esophageal motility, incomplete mastication of solid food or diaphragmatic stricture. It is important to note that the human \(\alpha\)-skeletal actin promoter directs Dp116 expression in skeletal muscle throughout the length of the esophagus (Fig. 6B), and thus, esophageal pathology cannot be attributed to loss of DGC expression in skeletal muscle.

**Dp116:mdx:utrn \(2/2\) mice display fragmented neuromuscular junctions**

The DGC also functions to promote the maturation and maintenance of the neuromuscular synapse (31,32). The neuromuscular

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**Figure 3.** Histology of quadriceps muscles from 8-week-old transgenic and control mice. (A) Entire cross-sections from the mid-belly of quadriceps muscles indicate extensive degeneration in both mdx:utrn \(2/2\) and transgenic Dp116:mdx:utrn \(2/2\) muscles compared with mdx. Scale bar: 1 mm. (B) High power (20×) fields of the muscles shown above along with wild-type control. Pathology of mdx and mdx:utrn \(2/2\) skeletal muscles is qualitatively similar at this age, with areas of extensive necrosis, inflammatory infiltrate and large numbers of myofiber containing centrally located nuclei. Similar pathologic changes are seen in the Dp116 transgenic muscles. Scale bar: 50 \(\mu\)m.
junction in muscles of \textit{mdx:utrn}^-/- mice is highly fragmented, with a reduced number of synaptic folds (9,29). This could simply be a consequence of muscle degeneration, or a direct result of reduction or absence of DGC members from the neuromuscular synapse. To test this possibility, we analyzed synapses in wild-type, \textit{mdx:utrn}^-/- and \textit{Dp116:mdx:utrn}^-/- mice. We found that the Dp116 transgene had no effect in preventing synaptic fragmentation or maintaining the normal number of synaptic folds (Fig. 7). In contrast, the \Delta\text{cys} dystrophin mutant protein (Fig. 1A) has been reported to reduce the degree of fragmentation of the neuromuscular synapse despite lacking a functional dystroglycan-binding domain (29). Here we show
that restoration of the C-terminal region of dystrophin and the DGC had little effect on synapse maturation and maintenance in \textit{mdx:utrn} \textsuperscript{2/2} mice. These results are consistent with previous assumptions that synapse fragmentation is caused by muscle degeneration (33).

\textbf{Dp116 does not confer mechanical stability to myofibers}

To decipher the mechanism through which Dp116 contributes to phenotypic improvement in \textit{mdx:utrn} \textsuperscript{2/2} mice, we systemically injected young \textit{mdx:utrn} \textsuperscript{−/−} mice with a recombinant adeno-associated viral vector (rAAV6) carrying the Dp116 transgene. rAAV6 is a small, non-integrating vector that is known to efficiently transduce mature skeletal muscle (34). At 4 weeks post-injection, we observed robust and widespread Dp116 expression in multiple muscles (Fig. 8). However, treated mice failed to demonstrate any overt phenotypic improvement when compared with untreated animals and continued to display growth retardation and contractures (data not shown). Immunofluorescent analysis revealed that Dp116-positive fibers were gradually lost from skeletal muscle following the initial peak in expression at 4 weeks. By 10 weeks post-injection, Dp116 expression in skeletal muscle was severely reduced or absent, indicating that episomal rAAV6-Dp116 genomes were lost during myofiber necrosis and regeneration. In contrast, Dp116 expression was retained for up to 5 months post-injection in \textit{mdx} muscles, which exhibit a milder dystrophy compared with \textit{mdx:utrn} \textsuperscript{−/−} muscles (data not shown). Thus, it is apparent that the loss of rAAV6-delivered Dp116 is dependent on the severity of dystrophic turnover, rather than any inherent instability of the construct. The loss of vector during muscle regeneration results in part from the inability of rAAV6 vectors to transduce muscle satellite cells \textit{in vivo}, such that muscle regeneration is not accompanied by replenishment of rAAV vector genomes from the muscle stem cell pool (Arnett \textit{et al.}, manuscript in preparation). These results demonstrate that Dp116-positive fibers are not mechanically protected from dystrophic turnover and implicate a non-mechanical mechanism for Dp116-mediated rescue in transgenic \textit{Dp116:mdx:utrn} \textsuperscript{2/2} mice.

\textbf{DISCUSSION}

Several transgenic and viral vector treated mouse models have previously succeeded in ameliorating \textit{mdx:utrn} \textsuperscript{−/−} muscle pathology to varying degrees by expression of different...
dystrophin and utrophin isoforms, as well as α7-integrin
(27,29,35–38). Conversely, transgenic mice expressing the
dystrophin isoform Dp71 or a dystrophin mutant with a dele-
tion in the cysteine-rich domain (Dcys) failed to improve the
phenotype of mdx:utrn\(^2/2\) mice (29,39). Although Dp71
does assemble the DGC, unlike Dp116 it has an incomplete
WW domain (Fig. 1A). Both the cysteine-rich region and
the WW domain are essential for dystrophin binding to
β-dystroglycan (40). The dystrophin–dystroglycan interaction
forms the central axis of the DGC and is likely to be in
dynamic equilibrium with other protein–protein interactions
(17,20,41–43).

Multiple groups have proposed that perturbation of signaling
pathways is an important mechanism of cell death in muscular
dystrophies caused by defects in the DGC (reviewed in 22,44).
These pathways could regulate a multitude of processes critical
for the maintenance of healthy muscle tissue, including apo-
tosis, hypertrophy, regeneration and vascular blood flow.
However, targeting of specific signaling pathways for potential
treatment of DMD has had limited success (45,46). In fact, no
specific signaling defect has yet been identified as a major
factor contributing to growth retardation in dystrophin-deficient
animal models, although the loss of nNOS leads to
exercise-induced ischemia and fatigue (47–49). One report
compared gene expression patterns of mdx and mdx:utrn\(^-/-\)
limb and extraocular muscles by DNA microarray (30). These
authors hoped to identify signaling pathways responsible for
the differential degree of pathology between genotypes and
between muscle groups that would manifest as alterations in
levels of gene expression. Interestingly, the most significantly
upregulated genes in mdx:utrn\(^-/-\) muscles were not known
signaling molecules but were proteins associated with the
slow muscle fiber phenotype. The authors proposed that this
expression pattern and fiber-type composition contributes to
the severe phenotype of mdx:utrn\(^-/-\) but may
instead be a consequence of, or a compensatory response to,
the cycles of myofiber degeneration and regeneration (50,51).
It is of interest that neuromuscular junctions associated with a
DGC unable to bind actin lack junctional folds despite the
relatively normal muscle mass (Figs 2 and 7).

Figure 7. Dp116 has no effect on the maturation or maintenance of the neuromuscular synapse in mdx:utrn\(^-/-\) mice. (A) The acetylcholine receptors stained with α-bungarotoxin (αBTX) are continuous in wild-type muscles, but fragmented in mdx:utrn\(^-/-\) and Dp116:mdx:utrn\(^-/-\) muscles. Electron microscopy (EM) images of the neuromuscular synapse shows a reduction in synaptic folds in mdx:utrn\(^-/-\) and Dp116:mdx:utrn\(^-/-\) muscles compared with wild-type
muscles. (B) Quantitative analysis of the data presented in (A) showing that the percent of continuous synapses and the density of folds in the postsynaptic membrane are not increased by expression of Dp116 in mdx:utrn\(^-/-\) mice. Bars show the mean ± S.D (*P < 0.001 compared with wild-type mice). Scale bar for images stained with αBTX is 15 μm and for EM images is 1 μm.
Most studies of dystrophin structure–function have generated evidence favoring the importance of a mechanical role in preventing dystrophic injury. Dystrophin mutations that perturb either the actin-binding or the dystroglycan-binding domains of dystrophin compromise the ability of the protein to prevent dystrophic damage to the muscle, even if all functional domains are present via simultaneous expression of two different dystrophin mutants (23–25,39,52,53). These results emphasize the importance of maintaining the mechanical link between the extracellular matrix and the actin cytoskeleton. We reported that systemic delivery of a micro-dystrophin gene using a viral vector led to increased muscle mass and extended the lifespan of mdx:utrn\(^{2/2}\) mice (27). In that case, expression of a truncated dystrophin (similar in size to Dp116 but containing a functional N-terminal actin-binding domain) increased myofiber size and specific force while reducing signs of dystrophic injury; none of these parameters was altered by Dp116 expression. Surprisingly, expression of Dp116 increased muscle mass and extended lifespan to a similar degree as the micro-dystrophin despite a lack of protection from dystrophy. Furthermore, the rapid loss of rAAV6-delivered Dp116 in mdx:utrn\(^{2/2}\) muscle clearly demonstrates that Dp116 does not protect myofibers from dystrophic degeneration (Fig. 8). The high degree of central nucleation and the histopathological appearance of Dp116:mdx:utrn\(^{2/2}\) muscle (Fig. 4) lend further support to this conclusion. Recombinant AAV6 is known to persist within the cell in an episomal state, and cellular turnover has been associated with the loss of vector genomes in vivo (54). As non-integrated structures, episomal vector genomes would be lost during fiber necrosis, and degenerated fibers subsequently replaced by newly generated, Dp116-negative myofibers in non-transgenic animals. In contrast, muscle regeneration in Dp116 transgenic mdx:utrn\(^{2/2}\) mice results in maintenance of Dp116 expression in regenerated myofibers, which leads to a significant phenotypic amelioration.

We interpret the above data as evidence for the lack of a mechanical function of Dp116, and that the mechanical function of full-length dystrophin primarily serves to protect against dystrophic injury. We cannot exclude the possibility that Dp116 contributes to stability of the sarcolemma through alternate interactions between the DGC and the cytoskeleton. It is clear that the network of cytoskeletal proteins associated with costameres is extensive, and dystrophin has been shown recently to interact with microtubules and intermediate filaments, interactions that may well occur with Dp116 (55,56). This may explain the modest improvement seen in resistance to contraction-induced injury, as restoration of a normal costameric lattice may improve lateral force transmission to adjacent myofibers (16). However, our results with both dystrophin-deficient and dystrophin/utrophin double-knockout mice demonstrate that any such interactions made by the Dp116/DGC complex are ultimately incapable of functionally substituting for the direct actin-binding properties of dystrophin or utrophin. The seemingly paradoxical improvement in muscle mass and function implicates a distinct function of dystrophin and the DGC that has not been previously described.

One cellular mechanism that is likely to be defective in mdx:utrn\(^{2/2}\) mice is regeneration. Until late in their lifespan, mdx mice show efficient regeneration of skeletal muscle that maintains muscle mass despite ongoing myofiber necrosis (8,57). Regeneration of skeletal muscle is primarily dependent on satellite cells, the resident stem cell population (58). Maintenance of regeneration has been proposed to explain the relatively mild phenotype of mice in which dystroglycan was

![Figure 8. rAAV6-delivered Dp116 does not protect myofibers from dystrophic turnover. mdx:utrn\(^{2/2}\) mice were treated with rAAV6-Dp116 at 2 weeks of age. Quadriceps (bottom) and TA (top) were collected and analyzed for Dp116 expression via immunofluorescent staining. At 4 weeks post-injection (left), robust expression of Dp116 is observed. In contrast, Dp116 expression is nearly absent at 10 weeks post-injection (right).](https://academic.oup.com/hmg/article-abstract/20/24/4978/591553)
disrupted specifically in skeletal muscle. These mice were generated using a muscle creatine kinase promoter driven Cre-loxP system to delete the dystroglycan gene in skeletal myofibers (59). This cross resulted in the complete disruption of the DGC in mature myofibers, but not in satellite cells or regenerating fibers. The authors concluded that expression of dystroglycan in muscle precursor cells is critical for continued regeneration, which efficiently compensated for muscle damage in their mice.

Our study is the first demonstration of an important functional role for restoration of the DGC without a concomitant link to the actin cytoskeleton by dystrophin or utrophin. When contrasted with the failure of Dp71 to produce a similar improvement in phenotype, our data suggest that a strong interaction between dystrophin and dystroglycan may be necessary for regulation of vital functions of the DGC that is separate from the linkage between the actin cytoskeleton and sarcolemma created by full-length dystrophin. We suggest that this function may be a signaling mechanism that regulates muscle growth and/or regeneration.

MATERIALS AND METHODS

Animals

To generate Dp116:mdx:utrn−/− mice, male Dp116:mdx4cv mice hemizygous for the Dp116 transgene were mated with mdx:utrn−/− females (9). The starting Dp116:mdx4cv transgenic mice had been backcrossed against the mdx4cv strain for more than six generations and were described previously (25). The Dp116-positive mdx:utrn−/− male progeny of this first cross was mated for a second generation with mdx:utrn−/− females. The second cross produced six possible genotypes: mdx, mdx:utrn−/− and mdx:utrn−/−, each either positive or negative for the Dp116 transgene. This breeding scheme was carried out independently with two different lines of Dp116 transgenic mice (2197 and 2354). All mice were genotyped by polymerase chain reaction at weaning age for both the utrophen mutation (60) and the Dp116 transgene (25). Selected mice were genotyped for the mdx mutation by an amplification-resistant mutation system assay (61). As expected, all mice tested were positive for the mdx mutation. Mice were weighed weekly after weaning and assignment of genotype. Both standard mdx and mdx4cv animals were used as controls for histological and physiological experiments. No significant differences were noted between the two genotypes so the data were combined and referred to as simply mdx in all figures. All experimental protocols were approved by the Institutional Animal Care and Use Committee of the University of Washington.

Measurement of serum creatine kinase levels

Blood was drawn from 8- to 10-week-old live mice via retro-orbital bleeds. The whole blood was immediately centrifuged and the serum collected and stored at 4°C. Measurement of the creatine kinase level was performed within 24 h of collection using the Stanbio CK, Liqui-UV kit (Stanbio Laboratory). For each measurement, individual samples were run in triplicate and compared with control normal and abnormal serum (Stanbio Ser-T-Fy I and Stanbio Ser-T-Fy II) to verify consistent results.

rAAV vector production and delivery

Flag-tagged Dp116 was inserted into an rAAV6 expression plasmid containing the cytomegalovirus immediate early enhancer + promoter, flanking inverted terminal repeats, and an SV40 polyA sequence. All vectors were prepared as described previously, using a two-plasmid co-transfection method into human embryonic kidney-293 cells followed by heparin-column purification (34). Vector titer was evaluated via southern analysis. Mice (n = 5) were injected intravenously with 1012 vector genomes (diluted in 200 μl isotonic saline) at ~2 weeks of age. An additional cohort of mdx (n = 8) and mdx:utrn−/− (n = 3) animals received IM injections (under isoflurane anesthesia) at a dose of 1010 vector genomes in 25 μl isotonic saline.

Immunofluorescence

Selected tissues were dissected, embedded in OCT (Sakura Finetek USA) and frozen in liquid nitrogen-cooled isopentane. Cryosections of 7–10 μm thickness were blocked in KPBS (20 mM potassium phosphate pH 7.4, 150 mM sodium chloride) containing 0.3 mg/ml bovine serum albumin (BSA) and 1% Tween 20. The blocked sections were incubated with primary antibodies diluted in KPBS containing 0.2% gelatin (KPBS-G) and 2% normal goat serum. Rabbit polyclonal antibodies were used to label Flag (Sigma), dystrophin NH2-terminus (53) and utrophin A (S. Froehner, unpublished data). After several washes with KBPS-G, the sections were incubated with goat anti-rabbit Alexafluor 488 (Molecular Probes). Immunostained slides were washed repeatedly with KBPS-G and mounted with Vectashield (Vector Laboratories). All photomicrographs were obtained with a Spot II CCD camera (Diagnostic Instruments, Inc.) and Spot Advanced software connected to a Nikon Eclipse E1000 using a ×20 Plan-Apochromat objective (numerical aperture = 0.75). Images for all the sections probed with a given antibody were acquired under identical exposure conditions.

Histological analysis and quantitative measurements of muscle fibers

Muscles were dissected, embedded in OCT (Sakura Finetek USA) and frozen in liquid nitrogen-cooled isopentane. Cryosections of 10 μm thickness were briefly fixed in methanol and stained with Gill’s hematoxylin and eosin-phloxine. The sections were washed, dehydrated and cleared in xylene before mounting with Permount. NADH staining was performed by incubating unfixed sections in 0.2 m Tris pH 7.4, 1.5 mM β-NADH reduced form disodium salt (Sigma) and 1.5 mM nitro-blue tetrazolium (Sigma) at 37°C for 30 min. The stained sections were washed for 2 min each in 30, 60, 90, 60 and 30% acetone and mounted in Vectashield (Vector Laboratories). Photomicrographs were obtained with a QCAM Fast cooled digital CCD camera and QCapture Pro software (QImaging) connected to a Nikon Eclipse E1000 microscope using a ×20 Plan-Apochromat objective.
Electron microscopy

Small (~1 mm³) regions of diaphragm muscles from 3-month-old wild-type, mdx:utrn −/− and mdx:utrn −/− mice expressing the Dp 116 transgene were fixed in half strength Karnovsky fixative for at least 24 h at 4 °C. Fixed muscles were washed in 1 × phosphate buffer (pH 7.1), post-fixed in 1% (w/v) aqueous osmium tetroxide containing 1.5% (w/v) potassium ferrocyanide (62), stained en bloc with uranyl acetate, dehydrated through acetone into Epon resin, embedded and polymerized at 60 °C for 24 h. Ultrathin sections were cut at ~ 80 nm, stained with Reynolds lead citrate (63), viewed and photographed with a Hitachi H600 transmission electron microscope (Tokyo, Japan). Synaptic folds were quantified using Image J (NIH) and compared using a one-way ANOVA with a Tukey post-test.

Neuromuscular synapse immunostaining

For wholemount preparations, skeletal muscle fibers were fixed for 2 h in 2% paraformaldehyde at 4 °C. Immediately following fixation, individual muscle fibers were teased apart using a fine needle. The fibers were then incubated 0.1 M glycine for 30 min, blocked in blocking solution (1% BSA, 0.05% Triton X-100 in 1 × PBS) for 1 h and stained with TRITC-conjugated α-bungarotoxin (αBTX; 1:500; Molecular Probes) for 1 h. The fibers were washed three times in blocking solution after staining and mounted on slides with antifade mounting media containing 4',6-diamidino-2-phenylindole (Vector Labs). Synapses were viewed and photographed using a Leica SL confocal microscope. Synapses were classified as continuous if they presented with three or less continuous regions of AChR clustering and discontinuous if they presented with more than three regions of AChR clustering. Approximately 600 synapses from n = 4 mice were analyzed and compared using a one-way ANOVA with a Tukey post-test.

Functional analyses

Force generation and susceptibility to contraction-induced injury was evaluated using a procedure previously developed by our laboratory (27). Briefly, male mice from 8 to 10 weeks of age were anesthetized with 2,2,2-tribromoethanol (Sigma) for in situ analysis of TA. Maximal force generation was evaluated at optimal muscle fiber length. Contractile performance was measured during a series of progressively increasing lengthening contractions stimulated at 10 s intervals. Following contractile evaluation, the TA was removed and weighed to calculate specific force. One-way ANOVA and Tukey post-test analyses were performed using GraphPad Prism software.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

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Conflict of Interest statement. None declared.

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