Exon skipping induces uniform dystrophin rescue with dose-dependent restoration of serum miRNA biomarkers and muscle biophysical properties

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Therapies that restore dystrophin expression are presumed to correct Duchenne muscular dystrophy (DMD), with antisense-mediated exon skipping being the leading approach. Here we aimed to determine whether exon skipping using a peptide-phosphorodiamidate morpholino oligonucleotide (PPMO) conjugate results in dose-dependent restoration of uniform dystrophin localization, together with correction of putative DMD serum and muscle biomarkers. Dystrophin-deficient mdx mice were treated with a PPMO (Pip9b2-PMO) designed to induce Dmd exon 23 skipping at single, ascending intravenous doses (3, 6, or 12 mg/kg) and sacrificed 2 weeks later. Dose-dependent exon skipping and dystrophin protein restoration were observed, with dystrophin uniformly distributed at the sarcolemma of corrected myofibers at all doses. Serum microRNA biomarkers (i.e., miR-1a-3p, miR-133a-3p, miR-206-3p, miR-483-3p) and creatinine kinase levels were restored toward wild-type levels after treatment in a dose-dependent manner. All biomarkers were strongly anti-correlated with both exon skipping level and dystrophin expression. Dystrophin rescue was also strongly positively correlated with muscle stiffness (i.e., Young’s modulus) as determined by atomic force microscopy (AFM) nanoindentation assay. These data demonstrate that PPMO-mediated exon skipping generates myofibers with uniform dystrophin expression and that both serum microRNA biomarkers and muscle AFM have potential utility as pharmacodynamic biomarkers of dystrophin restoration therapy in DMD.

INTRODUCTION
Duchenne muscular dystrophy (DMD) is an X-linked, monogenic muscle-wasting disorder and the most common inherited myopathy affecting children. DMD is caused by loss-of-function mutations in the gene encoding dystrophin (DMD), which performs structural and signaling functions at the sarcolemma.1–3 The leading approach for treating DMD is exon skipping, whereby splicing of the DMD pre-mRNA is modulated using antisense oligonucleotides, such that the translation reading frame is corrected, and an internally truncated, partially functional quasi-dystrophin protein is produced.4 At the time of writing (August 2022) there are four FDA-approved exon skipping drugs for DMD: eteplirsen, viltolarsen, golodirsen, and casimersen (all unconjugated phosphorodiamidate morpholino oligonucleotides [PMOs] designed for skipping of DMD exons 51, 53, 53, and 45, respectively).5 However, the levels of dystrophin achieved in treated DMD patient muscle are very low (~1% of healthy levels).5,6 Studies of dystrophin restoration therapies have primarily focused on the amount and quality of dystrophin protein expressed.5 In contrast, the importance of correct dystrophin localization has been less well studied. Notably, recent work from our group,7 and others,8 has suggested that uniform dystrophin expression is an important factor in the success of dystrophin restoration therapies and the severity of dystrophinopathies, respectively. Furthermore, challenges in determining dystrophin expression (e.g., the requirement for invasive biopsy, whether a single biopsy is reflective of the overall musculature, and technical difficulties associated with accurate measurement of dystrophin expression) have motivated the search for alternative DMD biomarkers.9 For example, extracellular micro-RNAs (ex-miRNAs) constitute one such class of minimally invasive biomarkers that may have utility in the context of DMD.10 The leading candidate miRNA biomarkers for DMD are the myomiRs (muscle-enriched miRNAs that regulate myoblast proliferation and...
In DMD, myomiRs (ex-myomiRs) offer excellent discrimination between DMD patients and healthy controls, suggesting they could be used to diagnose patients. However, the myomiRs have also been found to be elevated in the serum in multiple other neuromuscular conditions (including myotonic dystrophy, spinal muscular atrophy, amyotrophic lateral sclerosis), meaning that they are not specific biomarkers for DMD. Importantly, diagnosis of DMD is not currently a major clinical challenge. Instead, these ex-miRNAs may have utility as minimally invasive biomarkers for monitoring disease progression or response to therapy, currently unmet clinical need.

A single intravenous treatment of the mdx DMD mouse model with peptide-PMO (PPMO; designed to skip Dmd exon 23 and thereby rescue dystrophin expression) leads to restoration of ex-myomiRs toward wild-type levels. Similarly, ex-myomiR levels in dystrophic mouse serum are restored to near-wild-type levels after treatment with the U1 and U9 small nuclear RNA expressed exon skipping systems. Importantly, we observed that treatment with two PPMO conjugates with different potencies resulted in different degrees of ex-myomiR restoration, suggesting that ex-myomiR levels might be suitable for non-invasively assessing dystrophin levels in muscle. We further investigated this phenomenon using a genetic model (the mdx-Xist\(^{cShs}\) mouse) in which dystrophin is expressed at variable levels as a consequence of skewed X-chromosome inactivation. We generated animals that expressed ~10%, ~20%, and ~35% of healthy dystrophin protein levels, but surprisingly, ex-myomiR levels were not correlated with dystrophin protein expression at all, and were instead elevated at levels similar to those of dystrophin-null mdx controls, regardless of dystrophin expression. Importantly, dystrophin is expressed in a within-fiber patchy manner in the mdx-Xist\(^{cShs}\) model, suggesting that a uniform pattern of sarcosomal dystrophin is important for therapeutic correction. Here, we aimed to determine if PPMO-mediated exon skipping resulted in dose-dependent uniform dystrophin expression, ex-myomiR restoration, and correction of the biophysical properties of dystrophic muscle.

RESULTS

Dose-dependent dystrophin restoration in mdx muscle following PPMO treatment

Male mdx mice were treated with a single intravenous injection (via the tail vein) of a PPMO conjugate (Pip9b2-PMO) (Figures 1A and 1B) at 8 weeks of age. Mice were treated with one of three doses: 3 (n = 7), 6 (n = 9), or 12 mg/kg (n = 9). These treatments were intended to induce three different levels of dystrophin re-expression corresponding to each dose. Animals were sacrificed 2 weeks post treatment (i.e., at 10 weeks of age), at which point blood serum and tibialis anterior (TA) muscles were harvested. Untreated (age and sex matched) mdx (n = 9) and wild-type (WT; n = 5) animals were harvested in parallel as controls (Figure 1A). Dose-dependent restoration of dystrophin protein expression was confirmed by western blot with median expression values of 3.7%, 16.7%, and 44.8% of WT levels for the 3, 6, and 12 mg/kg PPMO-treatment groups, respectively (Figures 1C and 1D). Dystrophin protein was undetectable in untreated mdx TA muscle lysates (Figure 1C). The levels of exon-skipped (ΔExon23) Dmd transcripts in PPMO-treated muscle was similarly dose responsive (Figure 1E), as determined by RT-qPCR. Median skipped transcript levels were 1.2%, 11.8%, and 38% of total transcripts for the three PPMO-treatment dose groups, respectively. Dystrophin protein quantification and exon-skipped transcript levels were strongly correlated (Spearman’s r = 0.9262, p < 0.0001, Figure 1F).

Validation of myomiR RT-qPCR assay specificity

The leading serum miRNA biomarkers in the context of DMD are the myomiRs, consisting primarily of two miRNA families: (1) the miR-1/206 family (i.e., miR-1a-3p and miR-206-3p) and (2) the miR-133 family (i.e., miR-133a-3p and miR-133b-3p). Since that initial report, several groups have reported serum miRNA quantification data for these two miRNAs in the context of DMD, and human fetal myogenesis. These studies utilized the small-RNA TaqMan RT-qPCR method (from Applied Biosystems, now Thermo Fisher Scientific), which involves an RNA-specific stem-loop primer in the reverse transcription (RT) step. The small-RNA TaqMan approach is by far the most widely adopted miRNA quantification technology at use in this field (Figure S2B, Table S1). We were therefore motivated to determine if these assays are sufficiently specific to distinguish between miR-1a-3p and miR-206-3p, and between miR-133a-3p and miR-133b-3p. To this end, we generated artificial samples consisting of 50 fmol solutions of synthetic single-stranded RNA oligonucleotide mimics of each miRNA (Table S2, IDT) in nuclease-free water. Each artificial sample contained a single type of miRNA and was subsequently subjected to RT using either the correct RT primer that corresponded to the respective miRNA or the incorrect RT primer that corresponded to the related off-target miRNA. The resulting cDNA was analyzed in separate reactions using both the on-target and the off-target qPCR.
Figure 1. Dose-dependent dystrophin restoration in *mdx* mice treated with PPMO conjugates

(A) Schematic of experimental design. Eight-week-old male *mdx* mice were treated with a single intravenous injection of PPMO conjugate (at a dose of 3, 6, or 12 mg/kg) and harvested for analysis 2 weeks later. (B) Structure of the PPMO conjugate, which consists of a Pip9b2 peptide covalently conjugated to a PMO antisense oligonucleotide designed to skip *Dmd* exon 23. (C) Western blot of dystrophin protein expression for all PPMO-treated *mdx* TA muscle lysates. Vinculin was used as a loading control. (D) Tukey boxplot of dystrophin protein expression quantification. Statistical significance was tested by one-way ANOVA and Bonferroni post hoc test. *p < 0.05, **p < 0.01, ****p < 0.0001. (F) Correlation plot of dystrophin protein expression and exon 23 skipping levels for PPMO-treated *mdx* samples. Values were analyzed using linear regression and Spearman correlation analyses.
Figure 2. Stem-loop primed reverse transcription does not discriminate between miR-133a-3p and miR-133b-3p

(A) Sequence alignments for the miR-1/206 and miR-133 myomiR families. Differences between miRNA family members are highlighted in yellow. Artificial samples containing 50 fmol of synthetic miRNA mimic oligonucleotide templates were amplified using either the intended on-target small-RNA TaqMan assay (i.e., both RT primer and qPCR primer/probe mixes) or the assay for the related off-target miRNA. Amplification plots are shown for (B) miR-1a-3p and miR-206-3p and (C) miR-133a-3p and miR-133b-3p (the threshold is indicated by a dotted line). Diagrams of stem-loop reverse transcription primers in complex with either (D) their intended cognate target miRNA or (E) the unintended off-target related miRNA. High complementarity in the miRNA binding region for the miR-133 family stem-loop primers explains the off-target priming observed for miR-133a-3p and miR-133b-3p. FC, fold change.
assays. These experiments were designed to determine if (1) the RT primers can distinguish between closely related RNA targets and (2) whether the qPCR assays can distinguish between closely related cDNA templates.

Initially, each artificial sample was reverse transcribed with either the on-target or the closely related off-target RT primer, and then qPCR was performed using the correct qPCR assay. For the miR-1/206 family, the assays were found to be highly specific, with greater than 40,000-fold difference (ΔCq > 15.4) between on-target and off-target amplification (Figure 2B). Cq values for the off-target amplification were ~31–35, suggestive of low or very low levels of amplification. Conversely, the miR-133 family assays were not capable of discriminating between miR-133a and miR-133b, as equivalent amplification was observed when the miR-133a and miR-133b oligonucleotides were analyzed with either assay (ΔCq = 0.3) (Figure 2C). This lack of specificity could be attributed to the RT step, as negligible amplification was observed when the miRNA mimics were first reverse transcribed using the correct stem-loop primer, followed by qPCR with the off-target TaqMan assay (Figure S2). The reason for the cross-reactivity of these assays becomes apparent when the interactions between the stem-loop primers and the on-target/off-target miRNAs are inspected (Figures 2D and 2E). Stem-loop primers recognize their cognate miRNAs through the binding of the six 3’-terminal nucleotides of the RT primer with six complementary 3’-terminal nucleotides of the miRNA, which permits priming of the RT enzyme (Figure 2D).

For off-target primer interactions for the miR-1/206 family miRNAs, there are either two or three mismatches between the RT primer and the miRNA, including at the most 3’-terminal nucleotide, which is unlikely to support priming (Figure 2D). Conversely, for the miR-133 family, there is only one mismatch or one wobble base pair between primer and miRNA for the off-target RT primer combinations, respectively. In particular, the most 3’-terminal nucleotide is complementary in both cases, which is expected to support priming and therefore explains the high levels of off-target miR-133 amplification (Figure 2E).

These data demonstrate that small-RNA TaqMan assays are unable to distinguish between miR-133a-3p and miR-133b-3p, and are unlikely to be able to discriminate between other small RNAs that differ only at the 3’ terminus in the general case. For this reason, we have chosen not to investigate either miR-133a-3p or miR-133b-3p separately using RT-qPCR in our studies. Instead, we report only miR-133a-3p data (while acknowledging here that this assay will also amplify the less abundant miR-133b-3p species). Notably, we have previously shown that both miR-133a-3p and miR-133b-3p are upregulated in dystrophic serum using small-RNA sequencing (which can distinguish between these miRNAs).18

**Dose-dependent ex-myomiR restoration in mdx serum following PPMO treatment**

We next sought to analyze the serum levels of putative DMD biomarker myomiRs (miR-1a-3p, miR-133a-3p, miR-206-3p, and miR-483-3p) in the PPMO-treated animals, compared with WT and mdx controls. Levels of serum CK were measured in parallel. RNA was extracted from serum samples and miRNA levels were determined by small-RNA TaqMan RT-qPCR. All miRNA biomarkers were elevated in mdx serum (Figure 3). The myomiRs were significantly up-regulated in mdx serum with median fold-change increases of 16–19.9– and 48-fold for miR-1a-3p, miR-133a-3p, and miR-206-3p, respectively (Figures 3A–3C). miR-483-3p was upregulated by 12.5-fold but did not reach statistical significance at the p < 0.05 level (Figure 3D). PPMO-treated mdx mice exhibited a dose-dependent shift in all miRNA biomarkers toward WT levels. Specifically, the animals in the 12 mg/kg group exhibited biomarker restoration that was close to (and not significantly different from) WT levels, especially in the cases of miR-1a-3p and miR-133a-3p. Conversely, the animals in the 6 mg/kg group exhibited biomarker levels that were intermediate between those of mdx and WT animals (Figures 3A–3D). Highly similar results were observed for serum CK (Figure 3E). Notably, the variation in CK measurements was less than that observed for the miRNA biomarkers, and consequently more intergroup statistically significant changes were detected.

**Ex-myomiR biomarkers are anti-correlated with dystrophin expression**

Correlation analyses were performed to compare either muscle dystrophin expression or exon skipping levels with matched biomarker levels for PPMO-treated animals (Figure 4). Serum biomarker levels were strongly negatively correlated with dystrophin protein expression: miR-1a-3p, r = −0.72; miR-133a-3p, r = −0.76; miR-206-3p, r = −0.64; miR-483-3p, r = −0.55; and CK, r = −0.75 (all p < 0.005) (Figures 4A–4E). Similarly, serum biomarker levels were also strongly negatively correlated with exon 23 skipping: miR-1a-3p, r = −0.67; miR-133a-3p, r = −0.75; miR-206-3p, r = −0.67; miR-483-3p, r = −0.64; and CK, r = −0.75 (all p < 0.0006) (Figures 4F–4I). Conversely, the serum biomarkers were positively correlated with one another (Figure 4K, r > 0.72, p < 5.2 × 10^{-5}). miR-133a-3p was the most strongly correlated biomarker compared with dystrophin protein (Figure 4B), with serum CK a close second (Figure 4E). In contrast, miR-483-3p was the most weakly correlated biomarker (Figures 4D and 4I).

**Dose-independent uniform dystrophin distribution after PPMO treatment**

We previously showed that dystrophin distribution is uniform after treatment with a single 12.5 mg/kg dose of Pip9b2–PMO in the mdx mouse. This is in contrast to the patchy pattern of dystrophin expression observed in mdx-Xist mice expressing equivalent total levels of dystrophin compared with that achieved with exon skipping. We were therefore motivated to assess whether a within-fiber uniform pattern of dystrophin expression is observed at all PPMO doses tested. To this end, we analyzed dystrophin expression by immunofluorescence staining in TA muscle sections in both transverse and longitudinal orientations (Figure 5). PPMO-treated mdx mice generally exhibited a uniform pattern of dystrophin expression. In the low-dose (3 mg/kg) group, some degree of between-fiber
patchiness was observed when sections were viewed in transverse orientation (Figure 5). Specifically, some fibers were dystrophin positive, whereas others were dystrophin negative. The positively stained fibers generally exhibited consistent staining around the circumference of the fiber, although the strength of the staining was also weak in some fibers. In contrast, within-fiber patchiness was not observed, as viewed in longitudinal orientation (Figure 5).

**Dose-dependent restoration of myofiber elasticity following PPMO treatment**

A subset of the mice described above was further analyzed by atomic force microscopy (AFM) nanoindentation assay to investigate the relationship between dystrophin restoration, circulating biomarker levels, and the biophysical properties of muscle. AFM measures the resistance of a sample to an external deforming force, such that its stiffness can be determined. To this end, TA muscle explants (all n = 4) from WT, mdx, and PPMO-treated mdx mice at the three doses, 3, 6, or 12 mg/kg, were assessed by AFM. Force versus displacement measurements were performed, and Young’s modulus (E, a measure of the stiffness of the biological sample in response to an applied load) was calculated as described previously. An indentation depth of 1,050 nm was utilized such that Young’s modulus measurements reflected the stiffness of the myofiber inner structure, deeper than the actin microfilaments directly underlying the sarcolemma.

The mean E values were 4.7 and 1.7 kPa for the WT and mdx mice, respectively (a 2.7-fold reduction in mdx muscle, p < 0.0001), indicative of markedly lower resistance of dystrophic muscle to reversible deformation (Figure 6A), consistent with previous observations. Treatment with PPMO resulted in a dose-dependent shift in the mean E values (Figure 6A). Young’s modulus values were not significantly different between mdx mice treated with PPMO at the lowest dose, i.e., 3 mg/kg, and untreated mdx mice (Figure 6A), whereas the 6 mg/kg PPMO treatment was sufficient to completely reverse the reduction in stiffness observed in mdx muscle (Figure 6A). Interestingly, treatment with 12 mg/kg PPMO resulted in an ~2-fold increase (p < 0.0001) in Young’s modulus (mean E = 9.2 kPa) relative to the WT control group, suggesting a higher resistance to the deformation compared with that in the control mice (Figure 6A). Very similar
results were observed using measurement depths of 250 and 550 nm (Figure S3).

Inspection of underlying $E$ value determinations revealed a Gaussian distribution in $mdx$ muscle with an $E$ maximum at 1–2 kPa, consistent with high myofiber elasticity (Figure 6B). WT $E$ values exhibited a bimodal distribution with maxima at 2–3 and 7–8 kPa (Figure 6B). The PPMO-treated $mdx$ samples showed a gradation of effect, whereby the distribution of $E$ values shifted from an $mdx$-like distribution in the 3 mg/kg group to a more WT-like distribution in the 12 mg/kg group. For the low-dose 3 mg/kg group, the data were fit to a Gaussian distribution similar to the $mdx$ samples, although a smaller population of measurements with $E$ values ~5–7 kPa was also observed, which did not fit the calculated distribution. The results obtained for the 6 and 12 mg/kg data were best fitted with the sum of two Gaussian distributions, although the 6 mg/kg data were monomodal and the 12 mg/kg data were bimodal. The distribution of $E$ values in the high-dose group resembled that of WT muscle, although the $E$ values were shifted to the right (with a maximum at 6–8 kPa) and there were a high proportion of stiffer measurements (including up to 14–15 kPa), indicating an increase in myofiber stiffness in the PPMO-treated samples relative to WT (Figure 6B). Mean Young’s modulus was strongly positively correlated with both dystrophin protein expression ($r = 0.909$, $p < 0.0001$) (Figure 6C) and exon 23 skipping ($r = 0.902$, $p < 0.01$) (Figure 6D) in PPMO-treated $mdx$ mice. Overall, these findings suggest that exon-skipping-mediated dystrophin rescue can restore the biophysical properties of dystrophic muscle.

**DISCUSSION**

Here we show that treatment of dystrophin-deficient $mdx$ mice with Pip9b2-PMO results in dose-dependent $Dmd$ exon 23 skipping, dystrophin rescue, restoration of circulating miRNA biomarker levels, and correction of muscle biophysical properties. Treatment of $mdx$ mice with PPMO conjugates resulted in a within-fiber uniform pattern of sarcolemmal dystrophin expression across dose levels (Figure 5), consistent with our previous results. This is in contrast to our observations in CRISPR-Cas9-treated muscle (see related article), whereby dystrophin was expressed in a patchy manner. These findings point to a potential advantage of antisense oligonucleotide-mediated dystrophin rescue over gene editing.

This study adds to the growing body of literature that supports the use of ex-miomyRs as minimally invasive biomarkers for use in DMD patients. Notably, the negative correlations between circulating miRNA levels and molecular correction of dystrophin (Figure 4) suggest that these miRNAs could be utilized as pharmacodynamic biomarkers to assess the effectiveness of experimental therapies in clinical trials or approved therapies in patients. These findings are consistent with observations from a previous study from our group whereby two PPMO conjugates with differing potencies were found to induce corresponding levels of biomarker restoration. Importantly, several lines of evidence indicate that ex-miomyRs are not biomarkers of dystrophin expression per se. First, the elevation of serum miomyR levels (i.e., miR-1, miR-133, and miR-206) has been reported in the context of multiple other myopathies, and so is not a unique feature of dystrophinopathy. Second, serum miomyR biomarkers have been shown to be restored to WT levels in DMD mouse models following experimental therapies that do not involve dystrophin restoration. Specifically, serum miR-206 was restored to WT levels in $mdx$ mice treated with a myostatin pathway blockade strategy (i.e., in the context of no dystrophin restoration), and serum miomyRs are also present at WT levels in $mdx$-Fiona mice (which lack dystrophin but overexpress the dystrophin paralog utrophin).

Instead, we have suggested that ex-miomyR levels are reflective of muscle turnover (i.e., myonecrosis and compensatory regeneration) based on several lines of evidence: (1) ex-miomyRs are elevated during perinatal myogenesis in both WT and $mdx$ animals, (2) miomyRs are progressively released by human and murine myoblast cultures undergoing differentiation, and (3) a biphasic pattern of ex-miomyR release was observed in $mdx$ mice following an acute exercise regimen, corresponding to an initial damage phase immediately after exercise and a secondary regenerative phase 5–7 days post exercise. Furthermore, there is a degree of selectivity in miomyR release, suggesting that passive leakage from damaged muscle is insufficient to explain their serum abundance levels. Work from our group in the $mdx$ mouse has shown that ex-miomyRs are primarily non-vesicular and are instead protected from nucleolytic degradation through interactions with serum proteins.

Analysis of the biophysical properties of muscle using an AFM nanoindentation assay revealed reduced resistance to mechanical deformation (i.e., stiffness) in dystrophic muscle (Figure 6), consistent with previous reports. PPMO treatment restored myofiber elastic properties in a dose-responsive manner, although the high PPMO dose increased stiffness beyond that observed in WT mice. The reason for this is unclear, but given that treated $mdx$ mice exhibit signs of muscle hypertrophy (Figure 5), the resulting increase in the concentration of protein components of the contractile apparatus and cytoskeletal proteins may contribute to this increase in myofiber stiffness. In addition, the myonuclei of $mdx$ myofibers are often arranged in centrally nucleated chains, indicative of recent regeneration events, which may also contribute to an increase in stiffness. These data provide further evidence that serum miRNA measurements can be used to provide information about the underlying muscle (patho) physiology. Similarly, AFM has been used to assess the elastic properties of DMD patient biopsy material, suggesting that this technique...
could be utilized for diagnostic purposes. The data contained in this study provide evidence that AFM might also be suitable for assessing the effectiveness of therapeutic interventions for DMD (Figure 6), although invasive biopsy collection is still required.

Notably, in this study, serum CK arguably performed better than the ex-myomiRs. CK is known to be highly variable in patients and affected by a number of confounding factors (e.g., exercise, age, race, and drugs such as statins), which may not have an impact on serum myomiR levels, as has been reported by others. An important consideration is that the measurement of serum miRNA biomarkers requires multiple processing steps (i.e., RNA extraction, RT, qPCR amplification), whereas serum CK can be measured directly by either ELISA or enzymatic activity assay. As such, serum miRNA measurements may be subject to a greater degree of technical variation, especially in a research laboratory setting.

In this study, we also aimed to address the suitability of the small-RNA TaqMan RT-qPCR method to distinguish between closely related myomiRs. This commercial technology has been extensively used in the study of miRNAs across many areas of biology and medicine, and was shown to provide excellent discrimination between members of the let-7 family of miRNAs (some of which differ by only a single nucleotide, which is not the 3′ terminus). However, our data clearly show that this product is incapable of distinguishing between miRNAs that differ only in their 3′-terminal nucleotide, such as miR-133a-3p and miR-133b-3p, and that the lack of discrimination occurs at the RT step (Figure 2). We note that a similar failure to distinguish between miR-133a and miR-133b was reported by Ikeda et al., although the exact method and procedures used are not clear. This technical point is important, given that multiple studies have reported expression data for both of these miRNAs using this technology. The present study emphasizes the importance of assay validation, and how it should not be assumed that commercial products are valid a priori.

In conclusion, we show that PPMO-mediated exon skipping results in dose-dependent dystrophin rescue and restoration of the biophysical properties of dystrophic muscle. Importantly, a within-fiber uniform pattern of dystrophin expression was observed at all dose levels. Serum myomiR levels were inversely correlated with dystrophin expression and myofiber stiffness, thereby providing support for their use as pharmacodynamic biomarkers in the context of DMD.

MATERIALS AND METHODS
Animal studies
All procedures were authorized by the UK Home Office (project license 30/2907) in accordance with the Animals (Scientific Procedures) Act 1986. Animals were housed under a 12:12 h light-dark cycle, with access to food and water ad libitum. Mice used were male WT C57 (C57BL/10ScSn) or dystrophic mdx (C57BL/10ScSn-Dmdmdx/J).

PPMO
PPMO conjugates consisted of a peptide (Pip9h2, Ac-RXRRBBR FQILYRBRXR-OH, where X is aminohexanoic acid and B is β-alanine) covalently attached to a PMO designed to induce skipping of Dmd exon 23 (Table S2). The peptide was synthesized by standard Fmoc solid-phase chemistry and conjugated to the PMO via the amine of the morpholine heterocycle at its 3′ terminus, as described
Conjugates were dissolved in sterile water and passed through 0.22 μm cellulose acetate filters. Male mdx mice were injected with a single dose of PPMO conjugate (3, 6, or 12 mg/kg) in an equal volume of 150 μL 0.9% sterile saline via the tail vein at 8 weeks of age. Animals were sacrificed 2 weeks later (at 10 weeks of age) by escalating CO2 concentration, and serum/muscle tissues were harvested. TA muscles were macrodissected and flash-frozen in isopentane cooled on dry ice. Muscles were mounted on corks using Tissue-Tek O.C.T. Compound (VWR, Lutterworth, UK), to enable cryosectioning, and stored at −80°C. Blood was harvested from the jugular vein and collected using Microvette CB 300 collection tubes (Sarstedt, Nümbrecht, Germany) as described previously. Blood was incubated at 4°C for at least 2 h to facilitate clotting, and then cellular blood was pelleted by centrifugation at 10,000 g for 5 min. The serum supernatant was transferred to a fresh microcentrifuge tube and samples were stored at −80°C until ready for analysis.

Western blot
TA muscles were homogenized in modified RIPA buffer (50 mM Tris [pH 8], 150 mM NaCl, 1% IGEPAL CA-630, 0.5% sodium deoxycholate, 10% SDS, and 1× Complete protease inhibitor cocktail [Merck, Fethal, UK]) using a Precellys 24 tissue homogenizer (Bertin Instruments, Montigny-le-Bretonneux, France) (four cycles at 5,000 rpm for 30 s). Twenty micrograms of protein was mixed with NuPAGE sample reducing agent and NuPAGE LDS sample buffer and run on a NuPAGE 3% to 8% Tris-acetate gel (all Thermo Fisher Scientific, Abingdon, UK). Protein was electrotransferred onto a 0.45 μm polyvinylidene difluoride (PVDF) membrane for 1 h at 30 V, followed by 1 h at 100 V, and blocked with Odyssey blocking buffer (LI-COR Biosciences, Cambridge, UK) for 1 h at room temperature. Membranes were incubated with primary antibodies (Table S3) overnight in Odyssey blocking buffer supplemented with 0.1% Tween 20 at 4°C. Membranes were washed three times with PBST and then incubated with horseradish peroxidase (HRP)-conjugated secondary antibodies (Table S3) in Odyssey blocking buffer supplemented with 0.1% Tween 20 at room temperature for 1 h. Chemiluminescent signal was detected using Clarity Western ECL substrate (Bio-Rad, Watford, UK).

Exon 23 skipping RT-qPCR
Tissues were homogenized using a Precellys homogenizer (Bertin Instruments) (two cycles at 5,500 rpm for 30 s), and RNA was extracted using TRIzol reagent (Thermo Fisher Scientific) according to the manufacturer’s instructions. cDNA was generated from 500 ng of total RNA using the High-Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific) according to the manufacturer’s instructions. The resulting cDNA was diluted 1:5 prior to analysis. cDNA was amplified using a StepOne Plus real-time PCR thermocycler with TaqMan Gene Expression Master Mix (both Thermo Fisher Scientific) using universal cycling conditions: 95°C for 10 min, followed by 45 cycles of 95°C for 15 s and 60°C for 1 min. All samples were analyzed in duplicate. Absolute quantification was performed by comparing samples with standard curves comprising serial dilutions of the full-length (unskipped) and Δ23 Dmd (skipped) DNA target templates (IDT, Leuven, Belgium). The degree of exon 23 skipping...
was determined by calculating the percentage of skipped transcripts relative to the total (i.e., skipped + unskipped). Sequences of the RT-qPCR primers and probes are shown in Table S4.

**Immunofluorescence**

Fresh-frozen TA muscles were mounted onto corks with Tissue-TEK O.C.T. Compound and cryosectioned (8 μm) in transverse and longitudinal orientations. Sections were transferred onto SuperFrost Plus microslides (VWR), left to dry for 10 min at room temperature, and stored at −80°C until ready for analysis. Slides were air-dried, soaked in PBS for 10 min at room temperature, and then blocked in PBS supplemented with 20% fetal calf serum (FCS; Thermo Fisher Scientific) and 20% normal goat serum (NGS; MP Biomedicals, Eschwege, Germany) for 2 h at room temperature. Subsequently, the slides were incubated with primary antibodies in PBS supplemented with 20% PCS and 20% NGG for 2 h at room temperature. After being washed three times with PBS, the slides were incubated with fluorophore-conjugated secondary antibodies in PBS for 1 h at room temperature. The slides were then three times with PBS and mounted using Dako fluorescence mounting medium (Agilent, Didcot, UK). Details of all antibodies are shown in Table S3.

**Serum biomarker analysis**

Serum miRNA analysis was performed as described previously. Briefly, RNA was extracted from 50 μL of blood serum using TRIzol LS (Thermo Fisher Scientific) according to the manufacturer’s instructions, with minor modifications. A synthetic spike-in control oligonucleotide (i.e., cel-miR-39, 2.5 fmol, Table S2) was added at the phenolic extraction phase and mixed thoroughly. In addition, 150 μL of nuclease-free water (Thermo Fisher Scientific) was added at the phenolic extraction phase to increase the volume of the aqueous phase after phase separation. RNA was precipitated using isopropanol, and 1 μL of RNase-free glycogen (Roche, Welwyn Garden City, UK) was added to each sample as an inert carrier to assist RNA recovery. Pellets were air-dried, and then the RNA was resuspended in 30 μL nuclease-free water, and the samples were incubated at 55°C for 10 min. Samples were stored at −80°C until ready for analysis. RT was performed with the TaqMan MicroRNA Reverse Transcription Kit (Thermo Fisher Scientific) according to the manufacturer’s instructions, using 10 μL of each RNA sample. RT reactions were performed with miRNA-specific stem-loop primers as appropriate. The resulting RT product was diluted 1:2 prior to analysis, and 2 μL of cDNA was added per 20 μL qPCR. cDNA was amplified as described above. Data were normalized to cel-miR-3959 and analyzed using the Pfaffl method. Details of small-RNA TaqMan assays are provided in Table S5.

Serum CK analysis was performed as a service at the MRC Harwell Institute (Oxford, UK).

**Atomic force microscopy nanoindentation assay**

Mouse muscles were stored under liquid nitrogen in DMEM supplemented with 10% DMSO following the protocol published elsewhere. One hour before AFM measurements, muscles were thawed at room temperature by placing them in a Petri dish filled with DMEM (Sigma-Aldrich, Burlington, MA, USA). Afterward, they were washed twice in DMEM (for 2 min each time). The muscles were next removed from the medium, and the bottom part was gently dried with filter paper, followed by gluing them onto a glass coverslip using two 0.5 μL droplets of cyanoacrylate placed at both extremities as described previously. The muscle sample was immediately immersed in DMEM, and AFM measurements were performed (no longer than 3 h per individual muscle).

Measurements of the mechanical properties of muscles were carried out using an atomic force microscope (Xe120, Park Systems, Korea) equipped with a liquid cell sitting on a piezoelectric scanner with an xy range of 100 × 100 μm. AFM worked in a force spectroscopy mode that allows recording an approach and retracting the AFM probe (i.e., cantilever) using a separate piezoelectric scanner with a z range of 25 μm. Cantilevers were silicon nitride cantilevers of the ORC-8 (Bruker) type characterized by a nominal resonant frequency of 18 kHz, open angle of 36°, and nominal spring constant of 0.1 N/m. The spring constant of the cantilevers was calibrated using the Sader method. It ranged from 0.101 to 0.110 N/m. Force curves, i.e., dependencies between cantilever deflection and relative sample position, were recorded to estimate the elastic properties of muscles. Force curves were recorded in 36 positions (a grid of 6 × 6 pixels was set). For each muscle sample, 28–31 maps were recorded (in total, 1,008 to 1,116 force curves were analyzed). Measurements were acquired at an approach and retract speed of 8 μm/s. Calibration curves (curves recorded on a stiff, non-deformable surface) were collected from the measurements of a silicon nitride rectangular chip glued at the muscle height level.

The contact mechanics model analyzes the relationship between the load force and the indentation. In the AFM, to obtain indentation values, force curves recorded on a stiff, non-deformable surface (here, glass coverslip) were subtracted from force curves collected on a muscle sample. The resulting force versus indentation curves were fitted with the Hertz-Sneddon contact mechanics assuming that the indenting AFM probe can be modeled as a cone. For such a probe, the relation between the load force $F$ and the resulting indentation depth $\delta$ is:

$$F = \frac{2\tan \theta}{\pi} \frac{E_{\text{muscle}}}{1 - v^2} \delta^2,$$

where $\theta$ is the open angle of the probing cone (here, 36°), $v$ is Poisson’s ratio defining the compressibility of the studied material (here, $v = 0.5$ because cells are incompressible), and $E_{\text{muscle}}$ is Young’s (elastic) modulus of the studied muscle sample. Young’s modulus was determined at three indentation depths, i.e., 250, 500, and 1,050 nm.

**Statistics**

Statistical analyses were performed using GraphPad Prism v.9 (GraphPad Software, La Jolla, CA, USA). Differences between groups
were compared using one-way ANOVA and Bonferroni post hoc test. Differences were considered significant at the p < 0.05 level. Other analyses performed in GraphPad were the Shapiro-Wilk and Kolmogorov-Smirnov tests for normality, linear regression, non-linear regression (Gaussian and sum of two Gaussian), and Spearman correlation analysis. Data were tested for outliers using Grubb’s test, after which one sample in the 12 mg/kg group (which exhibited extremely high biomarker levels) was removed from all analyses. Heatmaps were produced using TMeV (The Institute for Genomic Research, Rockville, MD, USA).

**Data availability statement**
All relevant data are included in the article. Raw data are available on request.

**SUPPLEMENTAL INFORMATION**
Supplemental information can be found online at https://doi.org/10.1016/j.omtn.2022.08.033.

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**AUTHOR CONTRIBUTIONS**
T.C.R., Y.L., and M.J.A.W. conceived the study. K.C., J.O., J.A., M.L., N.A., A.M.L.C-S., G.M., and Y.L. performed experiments and acquired data. T.C.R., Y.L., and M.J.A.W. supervised the work. K.C., J.O., and T.C.R. wrote the first draft of the manuscript. All authors contributed to the final draft of the manuscript.

**DECLARATION OF INTERESTS**
M.J.A.W. is a founder, shareholder, and consultant for PepGen Ltd, a biotech company that aims to commercialize PPMO technology.

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