Targeted Exon Skipping to Address “Leaky” Mutations in the Dystrophin Gene

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Protein-truncating mutations in the dystrophin gene lead to the progressive muscle wasting disorder Duchenne muscular dystrophy, whereas in-frame deletions typically manifest as the milder allelic condition, Becker muscular dystrophy. Antisense oligomer-induced exon skipping can modify dystrophin gene expression so that a disease-associated dystrophin pre-mRNA is processed into a Becker muscular dystrophy-like mature transcript. Despite genomic deletions that may encompass hundreds of kilobases of the gene, some dystrophin mutations appear “leaky”, and low levels of high molecular weight, and presumably semi-functional, dystrophin are produced. A likely causative mechanism is endogenous exon skipping, and Duchenne individuals with higher baseline levels of dystrophin may respond more efficiently to the administration of splice-switching antisense oligomers. We optimized excision of exons 8 and 9 in normal human myoblasts, and evaluated several oligomers in cells from eight Duchenne muscular dystrophy patients with deletions in a known “leaky” region of the dystrophin gene. Inter-patient variation in response to antisense oligomer induced skipping in vitro appeared minimal. We describe oligomers targeting exon 8, that unequivocally increase dystrophin above baseline in vitro, and propose that patients with leaky mutations are ideally suited for participation in antisense oligomer mediated splice-switching clinical studies.

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Introduction

Dystrophin is an essential component of the oligomeric protein complex that protects the sarcolemma from the shearing forces of muscle contraction and relaxation. Diminished expression and/or production of a faulty dystrophin causes the X-linked recessive allelic conditions Duchenne (DMD) and Becker muscular dystrophy (BMD). DMD is the most common form of childhood muscle wasting, with patients losing ambulation before 12 years of age and, without appropriate healthcare and support, the condition is typically fatal in the second decade of life.¹ In contrast, BMD presents with more variable phenotypes and may display a slower clinical progression, with varying levels of impairment² and in some cases, only minimal symptoms such as cramps on exercise and elevation of serum creatine kinase.

Consequently, dystrophinopathies can be viewed as a spectrum, with DMD, the most severe phenotype being caused by the almost or complete failure of the protein. BMD typically describes those patients with milder symptoms and increased life expectancy resulting from internally deleted dystrophin isoforms retaining partial function. The wide range of dystrophinopathy phenotypes and the positive correlation between dystrophin levels and the age at which independent mobility is lost³ suggests that even a modest increase in expression of a functional dystrophin isoform could confer substantial clinical benefit.

The majority of DMD causing mutations are exon deletions that disrupt the reading frame, with two deletion prone regions of the gene identified at the 5′ end (minor hotspot) and involving exons 45–55 (major hotspot).⁴ The frame-shifting deletions lead to premature translational termination and destabilize the gene transcript through the induction of nonsense-mediated decay, whereas BMD-causing mutations are typically in-frame deletions that have limited impact on the critical binding domains. These observations led to the reading frame rule that correlates the genotype/phenotype relationship seen in DMD and BMD.⁵ Although more than 90% of mutations described in the dystrophin gene abide by this rule, there are exceptions where a protein-truncating mutation manifests as a mild phenotype, or an in-frame deletion has severe consequences (for review see).⁶ Some of these exceptions to the reading frame rule can be explained by the catastrophic effects of the mutation on crucial functional domains, whereas others arise from unanticipated consequences to the processing of the dystrophin mRNA. The in-frame, genomic deletion of exon 5 has been reported to cause DMD, rather than BMD as expected. Subsequent RNA studies revealed that the deletion of exon 5 led to the loss of exon 6 from the mature mRNA, resulting in a frame-shift that is consistent with a DMD phenotype, indicating cross-talk between exons during processing.⁷ Conversely, Ginjaar et al.⁸ reported a nonsense mutation in exon 29 that induced variable levels of skipping of this in-frame exon in three boys with DMD, leading to mild, moderate, and severe phenotypes within one family.

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Individuals with deletions encompassing exons 3–7 can present with clinical severities ranging from BMD to DMD, and several mechanisms have been proposed that might moderate the predicted DMD phenotype in these cases; restoration of the reading frame by a second mutation, ribosomal frame-shifting, the use of alternative promoters, re-initiation of translation at an in-frame start codon in exon 9, and alternative splicing. We consider the latter possibility most likely, as transcripts capable of being translated into a BMD-like protein have been detected in dystrophic muscle containing another leaky mutation, the deletion of exon 45. Prior et al. identified naturally occurring transcripts missing exons 44 and 45 that could be translated into a functional dystrophin isoform. Induction of functional dystrophin after re-initiation of translation or the use of alternative promoters from this region of the dystrophin gene must be considered highly unlikely, since this would eliminate crucial actin binding domains.

Antisense oligomer (AO) mediated exon skipping during pre-mRNA processing to restore the reading frame or excise exons carrying stop codons is emerging as a promising therapy for DMD. This intervention can give rise to a mature mRNA that is translated into an internally truncated, but partially functional BMD-like dystrophin isoform. Proof-of-concept of this approach was reported after direct intra-muscular injections of splice-switching oligomers targeting exon 51 to restore the reading frame in a subset of DMD deletion patients, using either 2′-O-methyl (2OMe) modified bases on a phosphorothioate backbone, or a phosphorodiamidate morpholino oligomer. More recently, dystrophin expression was demonstrated in muscle after systemic delivery of both oligomers to patients with DMD with amenable mutations.

Clinical evaluation of the morpholino oligomer targeting exon 51 (AVI-4658, now termed Eteplirsen) revealed that, although there were clear differences in dystrophin levels before and after oligomer administration, not all participants receiving the same dosage responded comparably. With this in mind, it is possible that variations in responses to exon skipping oligomers will be more pronounced when these are applied to mutations that naturally exhibited a degree of “leakiness” and result in variable disease severity.

Here we present in vitro data showing that leaky mutations in the minor deletion hotspot at the 5’ end of the dystrophin gene transcript are highly amenable to AO-induced exon 8 and 9 skipping, with only modest inter-patient variation. Several splice-switching AOs were evaluated, and similar trends in exon 8(+9) skipping efficiency by all AOs tested were observed in the different patient derived cells. We have previously shown that directing splice-switching oligomers to exon 8 resulted in human or canine dystrophin transcripts missing both exons 8 and 9. Indeed, in early canine exon skipping studies, RT-PCR carried out using primers targeting exons 1–9 failed to detect any splice switching, but when amplification conditions were modified to include exons 1–10, robust exon 8 and 9 skipping was detected. Although there appears to be cross-talk between dystrophin exons 8 and 9 during pre-mRNA processing, this communication appears one way, as splice-switching oligomers directed to exon 9 induce specific removal of the target exon alone. In summary, we have designed AOs that induce efficient exon 8 (+9) skipping, and propose that these compounds should be considered for clinical evaluation to address deletions of dystrophin exons 3–7 and 5–7.

Results
Preliminary AO screening
The splice motif predictor program, exonic splicing enhancer (ESE) finder was used to predict putative ESEs in exon 8. Using this information, splice-switching AOs were designed to induce skipping of exon 8, targeting the 3′ acceptor splice site, intra-exonic domains and the 5′ donor splice site, and synthesized as 2OMe modified bases on a phosphorothioate backbone. These AOs were transfected into normal human myoblasts as cationic lipoplexes at 100, 200, and 400 nmol/l and subsequent RT-PCR analysis showed most oligomers induced pronounced excision of exons 8(+9). The loss of exon 9 does not disrupt the reading frame and is not considered contra-indicatory, and AOs targeting the 5′ end of the exon (i.e., splice acceptor and ESE motifs) induced the most efficient exon excision. These AOs also elicited substantial levels of exon 8(+9) skipping in myogenic cells from DMD patient 1 (DM1) with a deletion of exons 3–7, as shown in Figure 1d. Although, all oligomers induced efficient exon skipping at the concentrations tested, H8A(−6+24) appeared most effective in that minimal full-length dystrophin transcript product was observed. Different primer sets were used to amplify this region of the mRNA to minimize size differences between amplicons from normal and DMD gene transcripts.

Evaluation of 2OMe AOs in DMD myoblast (DM) cultures
DMD myoblasts from different patients, three carrying deletions of exons 3–7 and one carrying an exon 5–7 deletion were used to evaluate the four AOs previously shown to induce robust exon 8(+9) skipping in normal myogenic cells. Myoblasts from two of the patients harboring deletions of dystrophin exons 3–7, DM1 (Figure 2a) and DM3 (Figure 2b) appeared more amenable to induced exon skipping than DM4 patient cells, carrying a deletion of dystrophin exons 5–7. DM4 (Figure 2c) showed an overall lower response to all AOs but nevertheless maintained similar trends, with H8A(−6+24) inducing the most efficient exon removal, followed by H8A(+57+83). Pooled data from three transfections experiments in DM1, DM3, and DM4 cells is shown in Figure 2d. Preparations of DM2 patient cells were of variable myogenic quality, and generated less consistent data (data not shown).

Evaluation of 2OMe AOs in DMD fibroblast (DF) cultures
The same 2OMe AOs, as cationic lipoplexes (10–400 nmol/l), were transfected into Ad5.f50.AdApt.MyoD transformed fibroblasts from six unrelated DMD patients with deletions of exons 3–7 (Figure 3). In three separate experiments, patient fibroblasts DF2 and DF4 showed inconsistent exon skipping in response to all AOs tested (data not shown). DF1 showed a consistently poor but nevertheless dose-dependant response to all 2OMe oligomers over the range of concentrations tested (Figure 3a). DF3 (Figure 3b), DF5 (Figure 3c), and DF6 (Figure 3d), also responded in a dose-dependent manner to all AOs tested, with H8A(−6+24)
inducing the highest levels of exon 8(+9) skipping. Pooled data from four transfections of DF1, DF3, DF5, and DF6 is shown in Figure 3e. Although the overall level of exon skipping varied, a clear dose-dependant response was evident and H8A(−6+24) was again one of the most consistent splice-switching oligomers.

**Evaluation of phosphorodiamidate morpholino oligomers in DMD myogenic cells**

Three sequences were selected for synthesis as phosphorodiamidate morpholino oligomers (PMOs) and were transfected by nucleofection into myogenic cells from patients with DMD (DM1–4), over the concentration range of 10–100 nmol/l (Figure 4a). Each oligomer induced a dose response in each of the myogenic cell strains, with DM1 again showing the most consistent response to the oligomers. H8A(−6+24) and H8A(−6+18) generated similar levels of exon skipping, whereas H8A(+57+83) was consistently less effective than the other AOs at all transfection concentrations. Pooled data (n = 4) from treatment of DM1–4 is shown in Figure 4b.

**Dystrophin expression following PMO conjugated to the cell penetrating peptide K transfection**

Myogenic cell strains DM1–4 were transfected with oligomers H8A(−6+18), H8A(−6+24), and H8A(+57+83), prepared as PMO conjugated to the cell penetrating peptide K (PPMOKK) at a concentration of 2 μmol/l. Seven days after transfection, protein and RNA were extracted from the cells for western blot and RT-PCR analysis. Extracts from normal myogenic cells indicate wild-type levels of protein, and untreated patient cells were included to assess base line dystrophin arising from “leaky” expression. The western blot membrane was probed with NCL-DYS1 and NCL-Hamlet1 to detect dystrophin and dysferlin, respectively (Figure 5a). The latter was included as a high molecular weight, muscle-specific loading control. In each of the untreated samples, dystrophin was detected, ranging from 3–7% of normal dystrophin levels (Figure 5b). Robust dystrophin expression was induced in DM1 and DM2 cells, reaching levels of 30–60% of those in normal myogenic cells, normalized to dysferlin expression. All patient cells showed increased dystrophin expression, with...
DM2 showing the greatest increase, from 4% in untreated to 43–62% in treated cells, representing a 10–14 fold increase (Figure 5c). DM1 also showed a marked increase in dystrophin levels from 7% in untreated cells, to 31–40% after oligomer application. DM3 and DM4 showed more modest increases in dystrophin, reaching 17–24% and 12–14% of normal dystrophin levels respectively, after oligomer treatment (Figure 5b,c).

RT-PCR analysis showed that in each case, the target exons were removed efficiently by all of the oligomers tested, with perhaps H8A(−6+18) being marginally less effective as indicated by the residual full-length product (Figure 5d,e). H8A(−6+18) induced the highest protein levels in DM2 and DM3, and resulted in levels of expression comparable to H8A(−6+24) in DM1 and DM4.

Discussion

Exon 51 was selected as the first target for proof-of-concept human exon skipping trials, as this strategy is potentially applicable to about 1 in 10 patients with DMD. Depending upon the specific database interrogated, the next most common dystrophin deletion subgroups should respond to skipping of exons 45, 44, or 53 (~1 in 13–20 boys), whereas skipping exon 8(+9) would only benefit ~1 in 30 individuals with DMD. Another consideration when selecting dystrophin deletions that would have the reading frame restored by exon 51 skipping was that these mutations are considered to be absolute, resulting in very low, or undetectable dystrophin. This was indeed the case for over half of the participants in the Muscular Dystrophy Exon Skipping Consortium trial, but dystrophin analysis of untreated muscle revealed that several boys had 1–2% of normal dystrophin levels, and one had an estimated 5% of control levels. In the absence of data from pretreatment biopsies, such amounts of dystrophin are likely to confound interpretation of the dystrophin induced by targeted exon skipping. However, rigorous dystrophin quantitation in the Morpholino studies, before and after treatment, revealed unequivocal but variable increases in dystrophin expression in muscle from some of the trial participants. Two
participants responded to oligomer treatment by showing substantial increases in dystrophin expression (2–18% and 0.9–17%), whereas a third individual was found to have 7.7% dystrophin from a baseline of zero. Clinical application of exon 8 skipping should not be delayed because of the relatively small proportion of patients with DMD who are likely to benefit, but rather, exon 8 should be considered a priority, since deletions in this region of the dystrophin gene have shown a propensity for inconsistent correlation between the genotype and phenotype, and therefore may prove highly amenable to exon skipping therapy. Since exon 8 (and 9) skipping has been reported to occur naturally\(^22,23\) this region could be highly amenable to splice intervention. Furthermore, the presence of low levels of endogenous high molecular weight dystrophin, despite the frame-shifting deletions of up to 300 kb, should preclude an immune response to the oligomer-induced dystrophin.

**Design of AOs to excise exon 8(+9)**

The *in silico* prediction program ESE finder 3.0\(^{20,21}\) was used to identify domains in dystrophin exon 8 that potentially influence pre-mRNA processing. In the first screen, AOs were designed to span the acceptor splice site, three intra-exonic domains predicted to contain ESEs implicated in exon recognition, H8A(−6+18), H8A(+42+66), and H8A(+57+83), shown in purple (\(n = 4\)) (mean + SEM).

**Figure 3** Evaluation of 2′-O-methyl (2OMe) antisense oligomers (AOs) targeting dystrophin exon 8 in MyoD transformed Duchenne muscular dystrophy (DMD) fibroblasts. (a–d) RT-PCR analysis from unrelated DMD patient fibroblast strains, all carrying a deletion of dystrophin exons 3–7, transfected with the four 2OMe AO lipoplexes, targeted to exons 8 at 10–400 nmol/l. Full-length (FL) and oligomer induced amplicons are indicated. (e) Densitometry was used to estimate relative exons 8 and 9 skipping induced by H8A(−6+18), shown in blue, H8A(+6+24), shown in red, H8A(+42+66), shown in green, and H8A(+57+83), shown in purple (\(n = 4\)) (mean + SEM).
Exon Skipping for Leaky DMD Mutations

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DM1 − 6 + 24 − 6 + 18 + 57 + 83

Normal

Dystrophin

Dysferlin

DM1 DM2 DM3 DM4

DT

∆ 9

∆ 8 & 9

DM1 DM3 DM2 DM4

DT

∆ 9

∆ 8 & 9

Figure 4 Evaluation of phosphorodiamidate morpholino oligomers (PMOs), targeting dystrophin exon 8, in Duchenne muscular dystrophy (DMD) myoblasts. (a) RT-PCR products from four DMD myoblast strains (DM1 (Δ3–7), DM3 (Δ3–7), and DM4 (Δ5–7) myoblasts) transfected with three PMOs using nucleofection. Full-length and oligomer-induced products are shown. (b) Densitometry analysis (n = 4) showing relative exons 8 and 9 skipping induced by H8A(−6+18) (blue), H8A(−6+24) (red), H8A(+42+66) (green), and H8A(+57+83) (purple) (mean ± SEM).

Figure 5 Dystrophin protein expression in PPMOk treated Duchenne muscular dystrophy (DMD) patient myoblasts. (a) Western blot showing dystrophin and dysferlin expression in PPMOk transfected (2 μmol/l), untreated (UT) DMD myogenic cells, and untreated normal human primary myogenic cells. Full-length dystrophin is 427 kDa, Δ5–9 dystrophin is ~402 kDa and Δ3–9 dystrophin is ~395 kDa. Dystrophin and dysferlin (230 kDa) were revealed by NCL-DYS1 and NCL-Hamlet1, respectively using the Western Breeze detection system. (b) Densitometry was used to estimate dystrophin expression relative to that in normal human myogenic cells, normalized to dysferlin. (c) Densitometry analysis showing relative dystrophin expression, determined by western blotting, as compared with untreated DMD myoblasts (fold increase) and normalized to dysferlin expression on the same blot. (d) RT-PCR analysis of exons 8 and 9 skipping levels. Full-length and oligomer-induced products are shown. (e) Densitometry analysis showing relative exons 8 and 9 skipping induced by H8A(−6+18) (blue), H8A(−6+24) (red), and H8A(+57+83) (purple). Data from untreated cells is shown in orange.
Using the splice site analyser tool (ibis.tau.ac.il/ssat/SpliceSite), the acceptor and donor splice sites of exons 8 and 9 were unremarkable as compared with those of the flanking exons. However, dystrophin exon 9 skipping occurs frequently in the absence of any oligomer intervention, and has been reported to occur in mice, dogs, and humans. Exon 9 skipping alone is commonly detected in untreated DMD patient cells, and although this did not appear to be influenced by oligomers directed to exon 8, the propensity for spontaneous exon 9 excision may contribute to the high levels of dual exon skipping.

Another feature that distinguishes exon 8 from many other dystrophin exons is the length of the preceding intron 7, in excess of 110 kb. Intronic length does not necessarily reflect ease of excision from the mature mRNA, as introns 1 and 2 are in excess of 190 and 170 kb, respectively, and exon 2, (62 bp) is not efficiently dislodged from the dystrophin mRNA in normal myogenic cells (K. Greet, S.D. Wilton, unpublished data). In contrast, exon 3 (93 bp) was readily removed with oligomers targeting ESE motifs near the beginning of the exon, whereas targeting the exon 3 donor splice site failed to induce any skipping (unpublished data). Another “leaky” mutation is the deletion of exon 45, which is preceded by nearly 250 kb of noncoding intervening sequence. These massive introns contain microsatellite repeats, short interspersed elements, long interspersed elements, and transposable elements, and may contribute to the relatively high genetic instability of the dystrophin gene. Nevertheless, intron length alone is not the most important feature contributing to gene instability within the major deletion hotspot, as the small 6.6 kb intron 49 shows a higher density of deletion breakpoints than the 37-fold larger intron 44.

We concentrated on targeting splice-switching oligomers to two regions in exon 8, the acceptor splice site and a splice enhancer-rich intra-exonic domain in the first-third of the exon. Subtle changes in length and oligomer annealing site can influence AO efficiency, and hence, additional splice-switching oligomers were designed to target the annealing sites. H8A(−6+18) and H8A(+42+66), and AOs H8A(−6+24) and H8A(+57+83) proved to be effective. Although exon skipping was induced by oligomers annealing to the exon 8 donor splice site, we did not further pursue this target region. Our earlier studies targeting other human dystrophin exons found that the acceptor site and beginning of the exon are generally the better annealing sites for AO-induced exon excision.

Oligomers H8A(−6+18), H8A(−6+24), H8A(+57+83), and H8A(+42+66) were selected for more detailed evaluation as 2OMe AOs after transfection into MD patient myoblasts and MyoD transformed DMD patient fibroblasts. Again, H8A(−6+24) was generally the most effective splice-switching compound, although the other three oligomers were also very effective when evaluated using the 2OMe chemistry. H8A(−6+18), H8A(−6+24), and H8A(+57+83) were synthesized as PMOs and PPMOk for further evaluation. Interestingly, when evaluated as a PMO in an in vitro titration study, H8A(+57+83) was not as efficient at splice switching as H8A(−6+18) and H8A(−6+24). However, when compared as PPMOks, all three oligomers induced similar levels of exon excision and protein production, suggesting a threshold effect at the single transfection concentration tested. This emphasizes the importance of dose response assays to identify the most efficient splice-switching oligomers.

The in vitro data presented here reveal the sequence H8A(−6+24) synthesized as the 2OMe AO chemistry as a lead candidate sequence for clinical application for amenable DMD mutations (e.g., Δ3–7, Δ4–7, Δ5–7, and Δ6–7). Although 2OMe AOs are chemically modified and have greater nuclease resistance than naturally occurring nucleic acids, degradation does occur. It could be surmised that although shorter 2OMe AO sequences (e.g., H8A(−6+18)) do induce splice switching, the longer the original oligomer (H8A(−6+24)), the more effective the degradation products. Differences in splice-switching efficiencies between H8A(−6+18) and H8A(−6+24) were less obvious when these sequences were evaluated as the more stable PMOs. In addition, the extra 6 bases in H8A(−6+24) do not appear to increase the potential for off-target effects, as both target sequences produced a single potential hit with 50 and 68% coverage, respectively, when submitted to a BLAST search of human genomic and transcript sequences (data not shown). However, if PMO oligomer production costs are considered, the shorter H8A(−6+18) may be more cost effective for clinical application.

Although shorter oligomers are more economical to produce and have less potential for cross-annealing to homologous sequences, it would be counter productive to use the shorter “inexpensive” oligomers if they are significantly less effective as splice switching agents. For example, H8A(−6+14), when transfected into normal cells as a 2OMe AO, induced only minimal exon skipping at concentrations of 300 nmol/l or above (data not shown). Nevertheless, it is sometimes difficult to distinguish relative skipping efficiencies of suboptimal compounds transfected into patient DMD cells, unless tested at low concentrations to allow discrimination of the more potent compounds. Exclusion of dystrophin exons 8 and 9 in cells from a patient missing exons 3–7, generates in-frame transcripts that escape nonsense-mediated decay, and the induced transcript would therefore have a longer half-life than the out-of-frame transcript. This could lead to overestimation of exon skipping efficiency, and hence, it is important to evaluate AOs in both normal and patient cells, and over a range of transfection concentrations.

Although myogenic cells from the patients with DMD are greatly preferred for oligomer evaluation in vitro, there are limitations to using muscle-derived cells from the patients with DMD, in particular, the invasive nature of muscle biopsies unless the patient is undergoing elective surgery. In addition, we have observed considerable variation in the proliferative and myogenic capability of cells obtained from dystrophic muscle, particularly when the tissue shows advanced dystrophic pathology. Although myogenic cells from different patients with DMD showed some variation in exon skipping efficiency in this study, both at the RNA and protein levels, exon 8(+9) was generally excluded in more than 50% of the transcript products after transfection at the higher concentrations. It is not possible to establish if this variation is because of the nature of the mutation (i.e., deletion breakpoints), the influence of genetic background on splicing, or limitations of in vitro cell culture, such as variations in myogenic capacity.

One alternative to using muscle-derived patient myogenic cells is the propagation of dermal fibroblasts and forced myogenesis. Although the dermal cells can be readily obtained with very minor discomfort to the patient, we have found that in vitro studies using fibroblasts tend to produce...
variable results. Although these cells are useful for RNA studies where typically >30% exon skipping was induced at the higher transfection concentrations, analyzing protein in these cells is more challenging, presumably reflecting relatively inefficient myogenic transformation.

When undertaking western blotting, we attempt to load equivalent amounts of normal, untreated and treated patient cell extracts to allow estimation of the relative amounts of base line and induced dystrophin. Western blot analysis showed that untreated myogenic cells from the DMD patients produced from 3 to 7% of the levels of dystrophin found in normal cells, consistent with reports of exon 3–7 (and 5–7) deletions as being "leaky," and not strictly conforming to the genotype:phenotype correlations. Acknowledging the limitations of extrapolating in vitro data, these basal levels of dystrophin expression are in line with those expected in muscle from an individual with a severe BMD phenotype. Dystrophin levels of 3% or less than those in normal muscle are consistent with a DMD phenotype. Nicholson et al. reported that patients with DMD with low, but nevertheless detectable amounts of dystrophin became nonambulant some 18 months later than those who had no measurable dystrophin. Patients with dystrophin levels of 3–7% of those in normal muscle could be expected to show severe BMD or the intermediate form of muscular dystrophy, consistent with the phenotypes reported for patients with mutations in this region of the dystrophin gene.

The variable basal levels of dystrophin expression in these patients could prove problematic when evaluating AOs in clinical studies, unless baseline dystrophin expression is established before treatment is initiated. Recent data on extended administration of Eteplirsen, an exon 51 splice-switching oligomer, indicates that the duration of treatment may be more important in determining the level of dystrophin expression than the actual dosage (personal communication). The dose escalating study reported by Cirak et al. extended over 12 weeks, and over that time the increases in dystrophin expression are in line with those expected in muscle from these patients carrying deletions in the minor deletion hotspot. Although dystrophin was present in untreated cells, there was an unequivocal increase in dystrophin levels after the administration of oligomer. Therefore, analysis of basal dystrophin expression in DMD muscle, before oligomer treatment, should be considered and imperative in clinical studies. It remains to be determined if those DMD/severe BMD patients with higher basal dystrophin levels will be more responsive to oligomer mediated exon skipping. The detectable dystrophin protein in muscle from these patients, without treatment, reduces the possibility of an immune response to the induced dystrophin isoform. We propose H8A(−6+24) to be a strong candidate for the treatment of DMD patients with deletions of exons 3–7 and 5–7 and further suggest that the propensity for these patients to produce "leaky" dystrophin is a good reason to expedite exon 8 skipping clinical studies.

### Materials and methods

**AO design and synthesis.** Oligomers consisting of 2OMe modified bases on a phosphorothioate backbone were synthesized on an Expedite 8909 synthesizer (Applied Biosystems, Melbourne, Australia), as described by Adams et al. AO sequences are shown in Table 1, with nomenclature described by Mann et al. Oligomers, identified as efficient splice-switching agents, were then synthesized by Gene Tools.

| Table 1 | AOs used in the study. Synthesized as 2OMe AO, PMO, and PPMO<sub>k</sub> |
|---------|----------------------------------------------------------------------------|
| AO Target | Sequence (5′→3′) | Length | 2OMe | PMO | PPMO<sub>k</sub> |
| H8A(−10+10) | GUA UCA ACA UCU GUU AGC AC | 20 | Y | | |
| H8A(−10+20) | UGG AUA GGU GGU AUC AAC UUC UGG AAG CAC | 30 | Y | | |
| H8A(−7+15) | GAU AGG UGG UAU CAA CAU CUG U | 22 | Y | | |
| H8A(−7+18) | GAU AGG UGG UAU CAA CAU CUG UAG A | 25 | Y | | |
| H8A(−6+14) | GGG GGU AAC AUC AUC UGU AA | 20 | Y | | |
| H8A(−6+18) | GAU AGG UGG UAU CAA CAU CUG UAG A | 24 | Y | Y | Y |
| H8A(−6+24) | UAG UGUG GAU UGG UAU CAA CAU CUG UAG A | 30 | Y | Y | Y |
| H8A(−3+18) | GAU AGG UGG UAC CAA CAU CUG U | 22 | Y | | |
| H8A(−42+66) | AAA CUU GGA AGA UGU AUG AUA UGU A | 25 | Y | | |
| H8A(−53+83) | GCC UAC UUL UUG AAG CAA AAC UUG GAA | 27 | Y | Y | Y |
| H8A(−96+120) | GCC UUG GCA UAA CUA CUU CAA UUG ACC CUU G | 25 | Y | | |
| H8A(−134+158) | AUG UAA CUU AAA AUG UUC UUC UUU A | 25 | Y | | |
| H8D(−13−12) | UAC ACA CUU UAC CUU UAG AGA AUA G | 25 | Y | | |

2OMe, 2′-O-methyl; AO, antisense oligomers; PMO, phosphorodiamidate morpholino oligomers; PPMO<sub>k</sub>, PMO conjugated to the cell penetrating peptide K.
Cell culture and myogenic conversion of fibroblasts. Normal human myoblasts were prepared essentially as described by Rando et al. from de-identified muscle biopsies, obtained with informed consent, during elective surgery or following diagnostic muscle biopsies. Similarly donated after informed consent, de-identified fibroblasts were obtained from skin biopsies taken from the patients with DMD (Table 2). The use of human tissue has been approved by the University of Western Australia Human Ethics Committee (approval number RA/4/1/2295). Normal myoblasts were proliferated and differentiated as described previously by Harding et al. All cells were plated at 3 x 10^4 cells/well in 24 well plates that had been sequentially pre-treated for 1 hour with 50 μg/ml poly D-lysine (Sigma, Sydney, Australia) and 100 μg/ml Matrigel (BD Biosciences, Sydney, Australia).

Following fine dissection of the skin biopsies, fibroblasts were cultured in Dulbecco’s modified Eagle medium (Invitrogen, Melbourne, Australia) supplemented with 20% foetal calf serum (Serana, Bunbury, Australia), 1% GlutaMax-I (Gibco, Melbourne, Australia), 10 U/ml penicillin (Invitrogen), 10 mg/ml streptomycin (Invitrogen), and 250 ng/ml amphotericin B (Sigma). Fibroblasts were converted to myoblasts through forced myogenesis by transfection with a MyoD expressing adenovirus Ad5.f50.AdApt.MyoD (The Native Antigen Company, Oxford, UK), and then differentiated in low serum media. Briefly, patient fibroblasts were plated at 3 x 10^4 cells/well in 24 well plates that had been sequentially pre-treated for 1 hour with 50 μg/ml poly D-lysine (Sigma) and 100 μg/ml Matrigel (BD Biosciences). Twenty-four hours later, media was changed to DMEM supplemented with 5% horse serum (Serana, Bunbury, Australia), 1% GlutaMax-I (Gibco, Melbourne, Australia), 10 U/ml penicillin (Invitrogen), 10 mg/ml streptomycin (Invitrogen), and 1% glutamine (Gibco, Melbourne, Australia) supplemented with 20% foetal calf serum (Serana, Bunbury, Australia).

PMOs were transfected by nucleofection using the Amazix 4D-Nucleofector and the Cell Line P3-X kit (Lonza, Basel, Switzerland). Cells were trypsinized and counted as described above, and 8 x 10^4 cells per treatment group pelleted at 600 g and resuspended in 22 μl of solution P3. The PMO was added to the cuvette with 20 μl of the cell suspension, which was then placed in the Nucleofector and pulsed with program CM-132 according to the manufacturer’s instructions. Pre-warmed Roswell Park Memorial Institute medium, (80 μl) (Invitrogen) was added to each cuvette immediately, and the cells were allowed to recover for 10 minutes at 37 °C. The 100 μl of cell suspension was then transferred to a fresh 1.5 ml microfuge tube containing 900 μl of differentiation medium.

Table 2 Summary table of DMD mutations and patient cells used in this study. DMD patient muscle-derived cells are identified as DM, and dermal fibroblasts converted to the myogenic lineage by forced myogenesis are shown as DF, western blot carried out (WB)

| ID  | Phenotype | Cell type | RNA effect | Mutation | 20Me | PMO | PPMO | WB |
|-----|-----------|-----------|------------|----------|------|-----|------|----|
| DM1 | DMD       | Myoblast  | Δ3–7       | (c.94-?_649+?del) | Y    | Y   | Y    | Y  |
| DM2 | DMD       | Myoblast  | Δ3–7       | (c.94-?_649+?del) | Y    | Y   | Y    | Y  |
| DM3 | DMD       | Myoblast  | Δ3–7       | (c.94-?_649+?del) | Y    | Y   | Y    | Y  |
| DM4 | DMD       | Myoblast  | Δ5–7       | (c.265-?_649+?del) | Y    | Y   | Y    | Y  |
| DF1 | DMD       | Fibroblast| Δ3–7       | (c.94-?_649+?del) | Y    |     |      |    |
| DF2 | DMD       | Fibroblast| Δ3–7       | (c.94-?_649+?del) | Y    |     |      |    |
| DF3 | DMD       | Fibroblast| Δ3–7       | (c.94-?_649+?del) |       |      |      |    |
| DF4 | DMD       | Fibroblast| Δ3–7       | (c.94-?_649+?del) |       |      |      |    |
| DF5 | DMD       | Fibroblast| Δ3–7       | (c.94-?_649+?del) |       |      |      |    |
| DF6 | DMD       | Fibroblast| Δ3–7       | (c.94-?_649+?del) |       |      |      |    |

PMOs, 2-oxymethyl DMD Fibroblast; DMD, Duchenne muscular dystrophy; PMO, phosphorodiamidate morpholino oligomers; PPMO, PMO conjugated to the cell penetrating peptide K.
amplify exons 1–10 in samples from the patients with DMD, or 3–10 in samples from normal muscle. After 30 cycles (myoblasts) or 35 cycles (MyoD converted fibroblasts), a 1 μl aliquot was removed and subjected to nested PCR, amplifying exons 1–10 (DMD) or 4–10 (normal), for 30 cycles using AmpliTaQ Gold (Applied Biosystems). Details of PCR primers (Geneworks, Adelaide, Australia) are shown in Table 3. Gel analysis, imaging, and sequencing. PCR products were resolved on 2% agarose gels in TAE buffer and relative exon skipping efficiency estimated by densitometry of the full length and oligomer-induced PCR products on images captured by the Chemi-Smart 3000 system (Vilber Lourmat, Marne-la-Vallée, France), as described previously. The identities of induced transcripts were confirmed by band stab isolation, purification of templates using UltraClean spin columns (MoBio, Carlsbad, CA) and DNA sequencing using BigDye V3.1 terminator chemistry (Applied Biosystems) as per manufacturer’s instructions. RT-PCR product sizes are summarized in Table 4. Sequencing was conducted at the Lotterywest State Biomedical Facility Genomics (Perth, Australia). Densitometric analysis was conducted using Bio1D-software (Scientific Software Group, Provo, UT).

Western analysis of dystrophin expression. Western blots were performed using a protocol derived from Cooper et al. and Nicholson et al. Seven days after transfection, cells were harvested by scraping, and resuspension in treatment buffer (50 μl/4.5 mg wet pellet weight) consisting of 125 mmol/l Tris-HCl pH 6.8, 15% sodium dodecyl sulfate, 10% glycerol, 0.5 mmol/l phenylmethylsulfonyl fluoride, 50 mmol/l dithiothreitol, bromophenol blue (0.004% w/v), and a protease inhibitor cocktail (15 μl/500 μl of treatment buffer) (Sigma). Samples were vortexed briefly, sonicated for 1 second, and heated at 95 °C for 5 minutes. Western analysis of dystrophin expression was performed at 18 °C at 290 mA, in a transfer buffer (20% methanol, 10% glycerol, 0.02 mmol/l Tris-Bis/Glycine sodium dodecyl sulfate gradient gel at pH 8.8, with a 3% stacking gel, pH 6.8. Fractionated proteins were transferred to a FluorotransW PVDF membrane (Pall, Melbourne, Australia) overnight at 18 °C at 290 mA, in a transfer buffer without methanol. Dystrophin was detected with NCL-DYS1 monoclonal anti-dystrophin (Novocastra, Newcastle upon Tyne, UK) applied at a dilution of 1:400, and dysferlin was detected using NCL-Hamlet1 at a dilution 1:1500 for 2 hours at room temperature. Detection was performed using a Western Breeze kit as per the manufacturer's instructions (Invitrogen). Enhanced chemiluminescence reactions were detected directly by the Chemi-Smart 3000 gel documentation system (Vilber Lourmat), using Chemi-Capt software for image acquisition and Bio-1D software (Scientific Software Group) for image analysis.

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### Table 3

| Application       | Primer | Sequence (5′→3′) |
|-------------------|--------|------------------|
| Normal RT-PCR    | Exon 3F | GGA AGC AGC ATA TTG AGA ACC |
| Normal Nested    | Exon 4F | GGA TCC ACA AGA GGT CAT GGC |
| DMD RT-PCR       | Exon 1F(o) | CTT TCC CCC TAC AGG ACT CAG ATC |
| DMD Nested       | Exon 1F(i) | GGG AGG CAA TTA CTC TCG GAG |
| RT-PCR reverse   | Exon 10R | CTC TCC ATC AAT GAA CTG CC |
| Nested reverse   | Exon 10R | GAC TTG TCT TCA GGA GCT TC |

### Table 4

| Diagnosis | Full-length (bp) | ΔExon 9 (bp) | ΔExon 8 (bp) | ΔExons 8 and 9 (bp) |
|-----------|-----------------|-------------|-------------|---------------------|
| Normal    | 788             | 659p        | 606         | 477                 |
| DMD Δ3–7  | 601             | 472         | 419         | 290                 |
| DMD Δ5–7  | 772             | 643         | 590         | 461                 |

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