Evaluation of exon-skipping strategies for Duchenne muscular dystrophy utilizing dystrophin-deficient zebrafish

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Abstract

Duchenne muscular dystrophy (DMD) is a severe muscle wasting disease caused by mutations in the dystrophin gene. By utilizing antisense oligonucleotides, splicing of the dystrophin transcript can be altered so that exons harbouring a mutation are excluded from the mature mRNA. Although this approach has been shown to be effective to restore partially functional dystrophin protein, the level of dystrophin protein that is necessary to rescue a severe muscle pathology has not been addressed. As zebrafish dystrophin mutants (dmd) resemble the severe muscle pathology of human patients, we have utilized this model to evaluate exon skipping. Novel dmd mutations were identified to enable the design of phenotype rescue studies via morpholino administration. Correlation of induced exon-skipping efficiency and the level of phenotype rescue suggest that relatively robust levels of exon skipping are required to achieve significant therapeutic ameliorations and that pre-screening analysis of exon-skipping drugs in zebrafish may help to more accurately predict clinical trials for therapies of DMD.

Keywords: dystrophin • DMD • Duchenne muscular dystrophy • zebrafish • muscle • exon skipping

Introduction

Mutations in the dystrophin gene can lead to two forms of muscular dystrophy. Null mutations that cause complete loss of dystrophin function typically result in Duchenne muscular dystrophy (DMD), a lethal disease characterized by progressive muscle wasting. In-frame deletions that lead to internally deleted dystrophin protein generally result in Becker muscular dystrophy (BMD), which shows milder symptoms than DMD with a large variation in clinical severity.

The finding of patients with a mild form of BMD that have in-frame deletions of often large regions of the dystrophin gene, together with the existence of revertant, dystrophin positive fibres in many DMD patients [1] suggests that exclusion of exons harbouring mutations can result in largely functional dystrophin protein. Similarly, the functionality of an artificial mini-dystrophin harbouring deletion of exons 18 to 58 has been reported suggesting much of the dystrophin protein may be dispensable for its function [2]. Currently, clinical trials are underway that are based on administration of exon-skipping antisense oligonucleotides (AO) [3, 4]. In exon skipping, AOs are employed to sterically block exon recognition splice motifs located in the pre-mature transcript. Effective blocking leads to faulty transcript processing that does not splice targeted exons correctly, so that they are excluded from the mature mRNA, resulting in a shorter, but partly to largely functional dystrophin. Although this approach will only be useful for the treatment of a subset of patients, it is estimated that 83% of all DMD patients would theoretically benefit from single or double exon skipping [5]. The exon-skipping approach has shown much promise in recently finished clinical trials [3, 4]. However, these clinical trials were limited, as the AO was only administered locally into a single muscle and healthy volunteers could not be tested due to the possibility of adverse affects on dystrophin function. Exon skipping has also been performed in the mdx mouse and in the golden retriever muscular dystrophy (GRMD) dog [6, 7], and the efficacy of exon-skipping protocols in generating Dystrophin protein has been demonstrated.
in both these models. However, the mdx mouse does not resemble the severity of the human condition and a systematic and statistical evaluation of exon skipping in dogs is hampered by their inefficient breeding and phenotypic variety [8]. Therefore, the level of dystrophin function necessary to effect full functional rescue in the context of a severe dystrophin-deficient pathology remains unclear.

In contrast to the most commonly used model, the mdx mouse [9], dystrophin-deficient zebrafish, named dmd\textsuperscript{14222a} [10], closely resemble the human condition in severity, onset and specific symptoms [11]. The progressive increase in expression of dystrophin protein at non-junctional sites in zebrafish correlates well with a number of pathological similarities between dystrophin-deficient zebrafish and patients suffering from DMD. Dystrophin-deficient zebrafish larvae show extensive muscle degeneration accompanied by fibrosis, inflammatory response, activation of the muscle stem cell compartment and greater variation in myofibre cross-sectional areas [11]. The detection of dystrophic muscle by reduction in birefringence [12], a light effect of the fish muscle visible under polarized light, enables efficient usage of dmd mutants in statistically evaluated experiments. In addition, zebrafish as a model system provide valuable advantages as its embryos develop ex utero, are translucent, amenable for drug screening [13], highly manipulable and genetically tractable [14], suggesting that dystrophin-deficient zebrafish could be a clinically relevant context in which to evaluate efficacy of therapeutic strategies.

In this study, novel dystrophin-deficient zebrafish mutants were isolated and used to evaluate efficacy of phenotype rescue by administration of morpholinos. Exon-skipping protocols were established that deliver a graded restoration of dystrophin levels demonstrating that only robust levels of induced exon skipping lead to sufficient dystrophin function capable of rescuing the dmd pathology. This analysis therefore provides a benchmark for the level of exon skipping that is required to effect clinically relevant pathological amelioration in the face of a fully penetrant dystrophic phenotype.

Materials and methods

Zebrafish lines and maintenance

The dmd\textsuperscript{tm90c} and dmd\textsuperscript{14222a} mutant alleles were obtained from the Tübingen Stock Collection [12]. All mutant lines were maintained in the TU (Tübingen) zebrafish strain. Genotyping was accomplished via PCR followed by restriction digestion (Fig. S1). All animal experiments were approved by MAS/2009/05.

Non-complementation screen and allele identification

Forty-eight males were mutagenized with N-ethyl-N-nitrosourea (ENU) as described [15] with the alteration of using 0.004% Tricaine (Sigma-Aldrich, St. Louis, MO, USA) during treatment. Surviving fish were out-crossed to females. The resulting F1 generation was crossed to heterozygous dmd\textsuperscript{14222a} +/+ fish and the offspring analysed for their birefringence at 3 days after fertilization (dpf). This non-complementation approach resulted in the identification of two novel dystrophin alleles after a survey of 243 genomes. To stabilize their genetic background, founder animals were back-crossed to TU over six generations before further experiments were performed. To determine the mutation rate, DNA was isolated from all F1 males and 1704 kbp at 25 randomly chosen loci were sequenced. Identified mutations were confirmed in independent PCR, resulting in a mutation rate of 1 mutation every 107 kbp. Cryostat cross-sections were stained with Harris haematoxylin and eosin or immunostained as described [11].

Morpholino oligomer injections

Morpholino oligomers were designed and named as described [16] and ordered from Gene Tools LLC (Philomath, OR, USA). Sequences were: Z32A(−18 + 7): 5'-gattatctctaaagacagactattt-3', Z32E(+83 − 107): 5'-ttcacttgacctccagacactc-3' and Z32E(+133 − 157): 5'-tgactcgacagctctgccac-3', targeting exon 32 as indicated by the number in brackets. Zebrafish embryos were injected at the 1- to 2-cell stage into the yolk with 1.4 nl of morpholino solutions of different concentrations. Isogenic, un.injected embryos from the same clutches were analysed as controls.

RNA extraction and semi-quantitative RT-PCR

Total RNA was extracted from larvae at different stages with TriReagent (Sigma) and dystrophin cDNA was prepared with the SuperScript III First-Strand-Synthesis System (Invitrogen, Carlsbad, CA, USA) using the primer 34cDNA: 5'-ccaaatcacttggcataccctcc-3'. The cDNA was then subjected to PCR performed with Advantage-HF 2 Polymerase Mix (Clontech, Mountain View, CA, USA) using the primers 30For: 5'-gacatc-3' and 34Rev: 5'-ctgaacaccactgacc-3'. Sequences were confirmed by sequencing. To determine the relative abundance of different dystrophin transcripts, we performed semi-quantitative RT-PCR using wild-type (WT) dystrophin transcript (549 bp) as internal control. For each morpholino concentration described, the PCR was optimized to keep the number of PCR cycles to a minimum. Proportions of different amplified dystrophin transcripts were quantified by measuring the band intensities using ImageJ (densitometry).

For statistical analysis of the correlation of morpholino efficiency and the ratio of dystrophic larvae, three clutches of 80–100 eggs from independent breeding pairs were injected with the described morpholino concentrations and 3 dpf hatched larvae were analysed for their muscle birefringence. Two larvae per clutch, with a dystrophic phenotype when applicable, were pooled and used for RT-PCR based analysis of dystrophin transcripts. Data are represented as means ± S.E.M.

Results

Screen for novel dystrophin alleles

In order to examine the effectiveness of the zebrafish dmd mutant for evaluating exon-skipping protocols of DMD patients, we
determined its conservation in comparison to the human dystrophin gene. Alignment of the human (muscle specific Dp427m isoform) and zebrafish dystrophin shows that 57% of the human protein sequence is identical (84% conserved) to zebrafish dystrophin. Additionally, the exon–intron structure is highly conserved: with the exception of the first and the last exon, each human dystrophin exon has a corresponding exon in the zebrafish sequence and skipping of any exon induces the same frame shift in the zebrafish and the human dystrophin transcript (Fig. S2). Thus, the zebrafish model has the potential to examine the functional consequence of exon skipping in the context of a highly penetrant in vivo model of DMD.

In order to study the feasibility and characteristics of phenotype rescue by exon skipping, the targeted exon should not encode for a region essential for dystrophin function. Unfortunately, the original dmd ta222a allele that we initially characterized harbours a premature stop codon in an exon encoding for a N-terminal calponin homology domain that is believed to be essential for actin binding [10, 17] and the only other identified allele, sapcl100, carries a splice site mutation, which itself is unsuitable for testing exon-skipping protocols [18]. Thus, we sought to generate new dystrophin alleles that would fall within exons appropriate for the evaluation of exon skipping.

To identify novel dystrophin alleles, a non-complementation screen based on the dmd ta222a fish was performed. Adult male zebrafish were treated with ENU and subsequently crossed to WT female to generate F1 founders. The F1 founders were subsequently crossed to heterozygous dmd ta222a/+ fish. Resulting offspring were screened for their levels of birefringence at 3 dpf. Non-complementation of the dmd ta222a allele, indicated by a reduction in birefringence, identified 2 novel dmd mutants. In addition, the zebrafish mutant dmd tm90c (synonym: sapje, sap cl90c), previously identified in a large scale screen but yet to be molecularly characterized [12], was also confirmed not to complement the dmd ta222a allele (Fig. 1A–D). Whole mount immunohistochemistry with antibody against dystrophin indicated loss of dystrophin in all three mutants and each of the mutants possessed a similar level of phenotypic severity compared to the original (Fig. 1A'–D'). Subsequent sequencing of the genomic dystrophin coding region in the mutants and comparison with the WT sequence led to the discovery of premature stop codons within exons 21, 32 and 53 for the alleles named dmd pc1, dmd pc2 and dmd tm90c, respectively (Fig. 1B''–D''). The identified mutations are widely distributed across the repetitive spectrin repeat region of dystrophin, which is often deleted within BMD patients and is also largely dispensable for the functional rescue of the mdx mouse by mini-dystrophin [2]. Thus, the mutations evident in the novel dystrophin alleles dmd pc1, dmd pc2 and dmd tm90c lie within regions suitable for exon-skipping protocols (Fig. 1E).

**Dystrophin restoration in dmd homozygotes via exon skipping**

AO, designed to sterically hinder specific splicing signals, can be employed to alter the splicing of the dystrophin pre-mRNA so that
the targeted exon is excluded from the mature mRNA leading to an internally deleted, but partially to largely functional dystrophin protein. In such an exon-skipping approach, removal of the exon may neither disrupt the open reading frame nor remove essential regions of the protein. The skipping of exon 32, which is mutated in dmd pc2, satisfies these two criteria.

An phosphordiamidate morpholino oligomer, Z32A(–18/110017), was designed to target the 5′-end of exon 32. To avoid nonspecific morpholino effects at high concentrations, various concentrations of Z32A(–18/110017) were administered. By comparison of abnormal embryo rates versus skipping efficiency, the optimal concentration of Z32A(–18/110017) for injections was established to be 500 μM (data not shown), a concentration which induced a slight curve in the angle of the body axis and a small delay in development. Two days after delivery of Z32A(–18/110017) into WT embryos, RNA was isolated and RT-PCR across exons 30–34 performed. Although RT-PCR on uninjected embryos led to detection of only WT dystrophin transcript, RT-PCR after injection of Z32A(–18/110017) resulted in two additional transcripts: one corresponding to a larger transcript arising from insertion of intron 31 upstream and one corresponding to a shorter amplicon originating from simultaneous skipping of the two exons 32 and 33 (Fig. 2A and B). The latter amplicon represents dystrophin transcripts schematically depicted in (B). Interestingly, the proportion of induced exon-skipped transcripts is, with high significance, increased in 2-dpf-old dmd pc2/pc2 homozygotes compared to WT control embryos (82% ± 0.6% versus 45% ± 0.5%, P < 0.01, n = 3) suggesting that nonsense-mediated decay proportionally reduced the levels of un-skipped stop codon containing transcript. (G) Administration of Z32E(+/33+157) leads to exclusion of 78 bp from the 3′-end of exon 32 and administration of Z32E(+/33+107) induces inclusion of the intron located upstream of exon 32. Combination of these two morpholinos, however, leads to a robust skipping of exon 32, which is still detected 21 days after administration. Again, the proportion of morpholino-induced transcript is significantly higher in 2-dpf-old dmd pc2/pc2 homozygotes than WT embryos (93% ± 2% versus 66% ± 3%, P < 0.01, n = 3). (H) Comparison of the zebrafish cryptic splice site elicited by Z32E(+/133+157) with the corresponding human dystrophin sequence and the human 5′ splice consensus [24]. ** indicates P < 0.01.
transcripts was declining, they could still be detected 7 days after Z32A(-18 + 7) delivery (Fig. 2F).

To determine, if administration of Z32A(-18 + 7) leads to expression of dystrophin protein, whole mount immunohistochemistry with antibodies against dystrophin was performed. Three days after administration of Z32A(-18 + 7) the proportion of the exon 31–32 deletion transcript is 4.7% ± 0.5% of all dystrophin transcripts (n = 8; data are means ± S.E.M.) and, in contrast to uninjected dmdpc2pc2 homozygotes, faint dystrophin expression could be detected in Z32A(-18 + 7)-injected homzygous dmdpc2pc2 at the same stage (Fig. 2B and C).

Under polarized light the zebrafish muscle shows a birefringence effect that is caused by parallel thread-like myofibrils within muscle fibres (Fig. 3A). This birefringence is reduced in dystrophin-deficient zebrafish, as the fibre organization is lost due to detachment and retraction of fibres [10]. Therefore, the birefringence assay was employed to survey if the restored levels of dystrophin are sufficient to rescue the dmdpc2pc2 phenotype. Z32A(-18 + 7)-injected homozygotes, however, demonstrate a level of muscle loss comparable to that detected in untreated dmdpc2pc2 homozygotes (Fig. 3B”) and Z32E(+133+157) leads to the detection of mono-nucleated infiltrates and in (C”) to necrotic myofibre. The arrowhead in (C”) points to a gap where a necrotic myofibre has been replaced by mucus. Scale bar: 50 μm.

phenotype rescue of dmdpc2pc2 homozygotes via exon skipping

Given the surprising complexity of the response to the blocking of the 3’ or acceptor splice site of exon 32, additional mechanisms were sought to increase the efficacy of in-frame skipping of exon 32 in dmdpc2pc2 homozygous mutants. In addition to blocking 3’ acceptor or 5’ donor-splice sites, targeting exonic splice enhancers (ESE) located within exons has also reported to lead to exon-skipping events [19]. Therefore, the program “RESCUE-ESE” was used to predict ESE located in exon 32 of dystrophin. Based on the prediction, the morpholino Z32E(+133+157) was designed against an ESE positioned close to the 3’-end of exon 32. RT-PCR, performed in the same manner as described above, on WT embryos treated with Z32E(+133+157) led to the detection
of an additional dystrophin transcript corresponding to the deletion of bp +97 to +174 of exon 32, activating a cryptic splice site at 5’-GAG gugaaa-3’ (Fig. 2D, G and H). Therefore, a second morpholino, Z32E(+83 +107), was injected into the offspring with Z32E(+133+157) and Z32E(+83 +107), which was designed to target this region. Administration of Z32E(+83 +107) alone promotes inclusion of the intron upstream of exon 32. However, administration of the combined morpholinos Z32E(+133+157) and Z32E(+83 +107), in a concentration of 25 μM each, induced robust skipping of targeted exon 32 without non-specific morpholino effects. Similar to the result already demonstrated with Z32A(−18 +7) injection, the proportion of the morpholino-induced transcript in relation to the WT dystrophin transcript is significantly higher in dmdpc2pc2 homozygotes (93% ± 2%) compared to WT embryos (66% ± 3%; P < 0.01, n = 3) (Fig. 2G). Although 2 days after injection of dmdpc2pc2 homozygotes with the combination of Z32E(+133+157) and Z32E(+83 +107) robust skipping of exon 32 was observed, the amount of morpholino-induced dystrophin transcript declined with progressing age of injected embryos. However, skipping of exon 32 was still detected 21 days after injection, suggesting that the stability of the morpholino AO in vivo and its efficacy in inducing exon skipping 3 weeks after injection may provide an effective therapeutic window of treatment (Fig. 2G).

In order to survey the effect of Z32E(+133+157) and Z32E(+83+107) administration on dmdpc2pc2 homozygotes, morpholino, Z32E(+133+157) and Z32E(+83+107) was analysed on cross-sections stained with haematoxylin and eosin, which confirmed that in contrast to untreated dmdpc2pc2 homozygotes, no muscle loss or enhanced fibrosis could be found (Fig. 3B’ and D’).

Collectively, administration of two morpholinos, one against a splice enhancer and the other blocking a cryptic splice site, induced a high level of exon 32 skipping leading to restoration of a robust level of dystrophin protein, sufficient to fully rescue the phenotype of dmdpc2pc2 homozygotes.

**Determination of the threshold level of exon skipping required to effect phenotypic rescue in dystrophin-deficient zebrafish**

In contrast to administration of Z32E(−18 +7) that only leads to weak skipping of exons 31 and 32 and does not result in phenotype rescue, simultaneous administration of Z32E(+133+157) and Z32E(+83+107) shows a robust skipping effect leading to a full phenotype rescue. We therefore surveyed in more detail the level of skipping efficiency necessary to achieve dmd phenotype rescue.

For this purpose, we in-crossed dmdpc2pc2 carriers and injected the offspring with Z32E(+133+157) and Z32E(+83+107) combined at an equal concentration of 12 μM. First, to test the accuracy of the morpholino administration, dmdpc2pc2 larvae were identified by PCR and subsequently analysed for their induced skipping effect individually. As shown in Figure S3, the variation in the detected skipping effects was relatively low. Therefore, in subsequent experiments with morpholino concentrations varying from 0 to 12 μM, two dmdpc2pc2 homozygous larvae were combined and the proportion of exon 32 skipped transcript was analysed in relation to all dystrophin transcripts by RT-PCR. Also, progression of the dystrophic pathology after each morpholino administration was assessed by analysis of the muscle birefringence at 3 dpf (Fig. 4). The extremes of the injected concentrations, 0 and 12 μM, resulted in a Mendelian ratio of 24.6% ± 0.5% of affected larvae and no detectable dystrophic phenotype, respectively (data in mean ± S.E.M.). Intermediate concentrations, however, resulted in partial rescue of the birefringence in a highly dose-dependent manner as
manifested by intermediate phenotype ratios. Collected data suggest that a skipping efficiency of about 10% results in 10% of detected dystrophic fish, representing a reduction of the expected Mendelian ratio by about half. In contrast to this partial rescue, a skipping efficiency of about 30–40% seems to result in less than 3% of injected larvae with a detectable dystrophic pathology, revealing that a skipping efficiency of 30–40% is sufficient to evoke a near full rescue of the dystrophic phenotype. As described above, injection of Z32E(+133+157) and Z32E(+83+107) into WT embryos resulted in lower exon-skipping efficiencies. Therefore, skipping efficiencies measured in homozygous mutants were also analysed in WT embryos. Although exon skipping could not be detected at morpholino concentrations of 0 μM, 1 μM and 2 μM, injection of 12 μM of the morpholino combination resulted in 12% ± 1% skipping efficiency, 6 μM in 3.5% ± 0.2%, 5 μM in 2.8% ± 0.3%, 4 μM in 1.5% ± 0.2% and 3 μM in 0.6% ± 0.1% efficiency.

### Discussion

This report describes the efficacy of exon-skipping approaches in the dystrophin-deficient zebrafish mutant *dmd* that accurately resembles the severe dystrophic muscle phenotype of human DMD. Statistical and systematic evaluation of exon-skipping efficiency demonstrates that relatively robust levels of exon skipping are required to achieve a significant level of phenotype rescue in a severe dystrophic pathology *in vivo*. 

Recently, two promising clinical trials employing exon skipping of dystrophin have successfully been completed [3, 4], but some unanswered questions remain. Amongst the problems that have been encountered is the fact that individual AO cannot generally be tested for side-effects in healthy volunteers, only small biopsies can be taken that are not statistically evaluated, and it remains unclear to which level dystrophin function needs to be restored to significantly improve the condition of DMD patients. Exon skipping can be surveyed in tissue culture, however, the canine model, as well as our survey, shows a large discrepancy in the outcome of exon-skipping experiments undertaken *in vitro* and *in vivo* [20], pointing to the importance of establishing rescue protocols *in vivo*.

Our comparison of the exon–intron structure of the human and the zebrafish dystrophin gene shows a remarkable degree of conservation. With the exception of the first and the last exon, this level of conservation enables modelling of skipping of human dystrophin exons in the zebrafish *in vivo*. 

One of the novel *dmd* alleles that we identified in a genetic screen, *dmd*\textsuperscript{pc2}, carries a mutation in exon 32, deletion of which does not disrupt the open reading frame nor results in deletion of an essential region of the dystrophin protein. This leaves us in the unique position of testing exon skipping in a genetically tractable model of DMD *in vivo*. We have established the experimental paradigm that enables us to survey exon-skipping experiments in our *dmd*\textsuperscript{pc2} mutant, measure the necessary skipping efficiency to rescue the *dmd* phenotype, and histologically survey the outcome of the rescue. Our data suggest that about 10% to 20% of normal dystrophin levels lead to only partial restoration of muscle function, whereas levels of at least 30% to 40% are needed to significantly improve muscle function in the context of the zebrafish dystrophic condition. These results are comparable to studies that correlated the level of dystrophin, quantified by antibodies, to the severity of symptoms evident within a given patient. Although patients with less than 10% of the normal level of dystrophin are reported to show a severe clinical phenotype of DMD [21], patients with 15% or less display a moderate to severe DMD phenotype [22] and BMD patients with mild symptoms are found to have dystrophin levels above 30% [22, 23].

Our observations show a highly significant difference between exon-skipping protocols performed on siblings and on mutants, probably due to nonsense-mediated decay of mutant transcript that harbours a premature stop-codon. This effect has to be taken into account when skipping efficiencies in dystrophin-deficient and WT zebrafish are compared.

Interestingly, administration of the AO Z32E(+133+157) activated a cryptic splice site at 5’-GAGggaaaa-3’, which is highly conserved between zebrafish and human beings. This potential cryptic splice site should be taken into account when AO are designed against exon 32 in human patients. It also reveals a surprisingly high level of conservation of the genetic structure of the dystrophin gene across a large evolutionary distance. Furthermore, to our knowledge, this is also the first time two splice switching AOs, which singly either promoted intron inclusion or activated a cryptic splice site, have been combined to induce robust and specific targeted exon skipping, and suggests a new mechanistic basis to promote exon skipping of individual exons.

The quality of life of DMD sufferers is not considered in the *dmd* fish model, any treatment of DMD needs to be evaluated in the context of individual human patients, as the use of exon-skipping strategies that induce less than optimal skipping efficiency may well have a significant therapeutic benefit. Nonetheless, these results highlight the importance of testing exon-skipping protocols in an *in vitro* model that features a comparable mutant context to patients for which a clinical trial is designed, in this way allowing a more accurate design and prediction of the clinical trial. As demonstrated here, the zebrafish as an animal model has the advantage of rapid screening techniques to generate alleles specific for exon-skipping strategies targeting specific exons.

In summary, the *dmd* fish can effectively be used to evaluate proposed therapeutic strategies for the treatment of DMD. We reveal that exon skipping is an effective mechanism for restoring dystrophin function in the context of a fully penetrant dystrophic phenotype and propose that the zebrafish model will be a valuable regent in identifying, designing and evaluating future therapeutic strategies for DMD.
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Conflict of interest

The authors confirm that there are no conflicts of interest.

Supporting information

Additional Supporting Information may be found in the online version of this article:

Fig. S1 dmd allele identification. The following primer pairs amplify the dystrophin locus, characteristic for each allele, followed by restriction digestion with the designated enzyme. Resulting DNA band patterns are as depicted. dmdpc2: pc1F 5'-cttgctactcctacagt-3' and pc2R 5'-'tgtgaagggagttgacctgtaagagcacaagttt-3', DraI. dmdpc1: tm90cF 5'-cttcaaaaggcttttaagaaag-3' and tm90cR 5'-'tgtgaagggagttgacctgtaagagcacaagttt-3', Ddel. dmdtm90c: tm90cF 5'-cttgctactcctacagt-3' and pc2R 5'-'tgtgaagggagttgacctgtaagagcacaagttt-3', BsaI. dmdta222a: ta222aF 5'-cttcaaaaggcttttaagaaag-3' and ta222aR 5'-gttgaagggagttgacctgtaagagcacaagtt-3', DraI. dmd083c: tm90cF 5'-cttcaaaaggcttttaagaaag-3' and tm90cR 5'-'tgtgaagggagttgacctgtaagagcacaagttt-3', Ddel.

Fig. S2 Comparison of human and zebrafish dystrophin. Numbered boxes represent exons of the human dystrophin isoform Dp427m with amino acid numbers indicated. The conservation of each exon in comparison to the zebrafish dystrophin sequence is written inside each box. Blue boxes represent exons that have the same number of amino acids in human and zebrafish and yellow ones have a different number of amino acids, as indicated below each yellow box.

Fig. S3 Variation of exon skipping in individual larvae. At 3 dpf, the exon-skipping efficiency was analysed in individual larvae administered with an equal mixture of Z32E(+133+157) and Z32E(+83+107) at a concentration of 12 μM each. The variation in exon-skipping efficiency is relatively low, allowing pooling of two larvae for subsequent analysis in further experiments.

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