The importance of genetic diagnosis for Duchenne muscular dystrophy

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ABSTRACT

Duchenne muscular dystrophy (DMD) and Becker muscular dystrophy are caused by mutations in the dystrophin-encoding DMD gene. Large deletions and duplications are most common, but small mutations have been found as well. Having a correct diagnosis is important for family planning and providing proper care to patients according to published guidelines. With mutation-specific therapies under development for DMD, a correct diagnosis is now also important for assessing whether patients are eligible for treatments. This review discusses different mutations causing DMD, diagnostic techniques available for making a genetic diagnosis for children suspected of DMD and the importance of having a specific genetic diagnosis in the context of emerging genetic therapies for DMD.

INTRODUCTION

Mutations in the dystrophin-encoding DMD gene underlie Duchenne muscular dystrophy (DMD) and Becker muscular dystrophy (BMD), a severe (DMD) and milder (BMD) form of inherited, progressive muscle wasting.1,2 Normally, the dystrophin protein acts as a shock absorber during muscle fibre contraction by linking the actin of the contractile apparatus to the layer of connective tissue that surrounds each muscle fibre.3,4 In DMD, mutations abolish dystrophin function either by disrupting the reading frame or by generating a premature stop codon (figure 1A, B). As a result, the connection between the actin cytoskeleton and connective tissue is lost and muscle fibres are easily damaged during contraction, leading to chronic muscle damage, inflammation and eventually replacement of muscle fibres by fat and fibrotic tissue and thus loss of muscle function.5

By contrast, individuals with BMD generally have mutations that maintain the open reading frame, allowing the production of dystrophin proteins with an internal deletion or duplication that can connect actin to the connective tissue, and thus are partially functional (figure 1C). Consequently, individuals with BMD often show a later onset and a slower disease progression, although there is phenotypic variation, with some patients being diagnosed in childhood and others being diagnosed in midlife or later.5,6 Almost all DMD patients will develop cardiomyopathy,7 and there is a high risk patients with BMD will develop cardiomyopathy regardless of the skeletal muscle phenotype. The incidence of dystrophinopathies is ~1 in 4000 (1 in 5000 for DMD and 1 in 20 000 for BMD in male live births).5,8,9 Since the DMD gene is located on the X-chromosome, DMD and BMD present primarily in males and are maternally inherited.

Correct DNA diagnostic analysis is crucial for DMD and BMD patients since it is important for optimal care and family planning, but also provides information on eligibility for mutation-specific treatments.11 In this review, we will explain the different types of mutations that have been reported for the DMD gene, discuss genotype–phenotype correlations; outline the genetic diagnosis of children suspected of having DMD; and discuss the importance of a specific genetic diagnosis in the context of emerging genetic therapies. A glossary of terminology is provided in table 1.

MUTATIONS IN THE DMD GENE

The DMD gene is the largest known human gene, containing 79 exons spanning 2.2 Mb.5 The mutation rate is relatively high; in one in three cases, DMD is caused by a de novo mutation. As such, new cases will arise even with good prenatal diagnostic tools and family counselling for known cases. This high mutation rate also underlies the large variation of mutations that has been identified for patients with DMD (figure 2).6,12 Although the majority of patients have a deletion (~68%) or duplication (~11%) of one or more exons, small mutations are found as well (~20% of patients). These deletions and duplications can occur anywhere in the gene, but are concentrated between exons 45–55 and exons 2–10 for deletions and duplications, respectively.13 The outcome of deletion and duplication mutations can be twofold. If the number of nucleotides in the exons that are deleted or duplicated is divisible by 3, the reading frame will not be disrupted. This will allow translation of the mRNA into a dystrophin protein that, although slightly shorter or longer in the centre, contains the N-terminal and C-terminal domains crucial for connecting the actin cytoskeleton to the extracellular matrix. These dystrophins retain some degree of function and are found in patients with BMD. By contrast, when the number of nucleotides of the exons deleted or duplicated is not divisible by 3, the reading frame is shifted, leading to the incorporation of aberrant amino acids into the protein during translation. Often, an incorrect reading frame contains many stop codons, leading to premature termination of translation and production of a truncated protein. The resulting dystrophins lack the crucial domain that connects to the extracellular matrix and are therefore not functional and generally not stable. These mutations are associated with DMD.

There are several ways by which small mutations can interfere with dystrophin production (figure 2C). Small deletions or insertions can disrupt the reading frame at an exon level, which...
like larger out-of-frame deletions or duplications involving one or more exons leads to non-functional dystrophins. Point mutations can convert a codon for an amino acid into a stop codon, which will also result in premature termination of protein translation. Finally, point mutations or small deletions and/or insertions can disrupt a splice site. During the splicing process, introns are removed by the splicing machinery, for which recognition of the splice donor (first two nucleotides of an intron (GU)) and splice acceptor (last two nucleotides of an intron (AG)) is crucial. Generally, an exon for which the acceptor or donor splice site is mutated will no longer be recognised by the splicing machinery, leading to the exon being excluded from the mRNA. Thus, splice-site mutations generally cause a single-exon deletion at the mRNA level, which will lead to a partially functional or a non-functional dystrophin depending on whether the omitted exon was in-frame or out-of-frame, respectively.

In <1% of patients, other types of mutations are found; for example, deep intronic mutations can lead to an intronic region being recognised as an exon by the splicing machinery (cryptic or pseudo-exons) leading to its inclusion in the mRNA.\(^6\)\(^12\) Cryptic exons can disrupt the reading frame and/or will contain stop codons preventing the production of functional dystrophin. Missense mutations are rare in patients with DMD, and the few that have been reported are located in the cysteine-rich domain of dystrophin and abolish the binding to β-dystroglycan, which in turn connects to the extracellular matrix.\(^12\)\(^14\) Therefore, in effect these missense mutations are similar to other DMD-causing mutations because they also disrupt the connection between actin cytoskeleton and extracellular matrix. Finally, a handful of cases caused by translocations involving the DMD gene have also been reported. These mutations separate one portion of the gene from the other, preventing a full transcript being produced. Notably, translocation mutations involving the DMD gene will cause a DMD phenotype in females.\(^15\)\(^16\) Normally, carrier females will be born with 50% of dystrophin positive muscle fibres due to random X-inactivation of the chromosomes carrying the mutated or the normal DMD gene. With translocation mutations, however, cells where the translocated DMD gene is inactivated will not survive due to a lower dosage of the autosome involved in the translocation. Thus, the unaffected DMD gene will always be inactivated and no dystrophin can be produced.

**GENOTYPE–PHENOTYPE CORRELATIONS**

Close to 10% of genetic mutations do not follow the reading frame rule,\(^6\)\(^12\) that is, patients with in-frame mutations can present with DMD and patients with out-of-frame mutations can present with BMD. Some frequent exceptions to the reading frame rule are discussed below. However, for certain mutations both patients with DMD and patients with BMD have been reported. Furthermore, for patients with BMD and DMD, disease severity can vary for the same mutations, sometimes even within the same family and there is also variation in the extent in which the heart is affected.\(^6\)\(^7\)\(^17\)\(^19\) As such, most likely genetic modifiers play a role as well in determining disease severity. Discussing these is beyond the scope of this review, and we refer interested readers to a recent review paper by Vo and McNally.\(^20\)

Generally, the location or size of deletions or duplications that lead to out-of-frame mutations do not affect the clinical phenotype; if the reading frame is disrupted, the C-terminal part of the protein cannot be produced and the protein will not be functional. There are, however, three notable exceptions. First, frame-shifting or nonsense mutations before exon 8 can present with DMD in certain cases.\(^21\) This is due to the presence of an alternative translation initiation sites in exon 6 or exon 8 that are activated by some mutations (eg, a deletion of exon 2 or a stop mutation in exon 1 activate an alternative translation
initiation side in exon 6), but not others (eg, an exon 2 duplication). If exons 3–7 are deleted apparently, it varies between patients whether or not the exon 8 alternative translation initiation site is used. Second, patients with nonsense mutations in in-frame exons can present with BMD. This can be explained by the fact that the mutation disrupts sequences involved in exon recognition and, consequently, the exon is occasionally skipped, thus bypassing the nonsense mutation but maintaining the reading frame. Third, patients with DMD with mutations flanking exon 44 show a slightly milder phenotype (‘mild DMD’), with, on average, a later loss of ambulation and a slower decline in the distance walked in 6 min compared with other patients. The most likely explanation is low-level spontaneous skipping of exon 44. Indeed, this has been reported in cultured cells from patients with deletions flanking exon 44. This suggestion is further supported by the fact that the dystrophin levels in these patients are on average higher than for patients with DMD with other frame-shifting mutations.

For in-frame mutations, the location and size of mutations can influence disease severity to some extent. In-frame mutations that abolish the extracellular matrix-interacting domain (encoded by exons 64–70) or all actin binding domains (encoded by exons 2–10 and exons 32–45) will not result in functional proteins and are therefore associated with DMD. In-frame deletions affecting the first 10 exons delete the first two actin binding domains, but spare the third one encoded by exons 32–45, typically leading to a ‘severe BMD’ phenotype. Deletions in the hotspot region (exons 45–55) are generally associated with a milder disease presentation. Deletions between exons 10 and 40 are milder still and are sometimes only associated with cramps and myalgia and are sometimes found in asymptomatic individuals. Finally, X-linked dilated cardiomyopathy patients have mutations in the DMD gene, but present only with a cardiac phenotype in the absence of skeletal muscle problems. Here, dystrophin is produced in skeletal muscle, while it is not produced or not functional in heart. While most carriers do not present with symptoms, some ‘symptomatic carriers’ present with cramps, myalgia or even progressive muscle weakness. Notably, female carriers are prone to develop dilated cardiomyopathy as well.

**ESTABLISHING A GENETIC DIAGNOSIS**

Physicians should suspect DMD when young boys present with impaired muscle function, frequent falls and Gower’s sign. Delayed speech is also frequently observed in young boys with DMD, probably due to the absence of a brain isoform of dystrophin. Serum analysis will reveal the presence of elevated muscle enzymes due to leakage into the bloodstream (most notably creatine kinase (CK), but also transaminases such as aspartate transaminase and alanine transaminase). Upon these findings (delayed muscle function, speech delay and high CK), patients are generally referred to neuromuscular specialists, who request genetic analysis of the DMD gene to confirm whether the patient has DMD.

Given that deletions and duplications of one or more exons are found in the majority (70%) of patients, it is most cost-efficient and labour-efficient to check for these mutations first (see table 2 for the costs of DMD diagnosis in selected European countries). The multiplex ligation-dependent probe amplification (MLPA) analysis is the most reliable test to identify exactly which exons are involved in deletions or duplications. This approach uses sets of probes that can hybridise to each of...
the DMD exons. Each set of probes consists of two oligonucleotides that hybridise adjacent to each other to a given exon. Probes have a tail to allow PCR amplification and one of the probes contains a ‘stuffer sequence’, the length of which differs for each pair of probes. Upon hybridisation, a ligation step joins the two probes and a PCR will be performed, which can only amplify ligated probes (ie, probes that were able to hybridise). The resulting PCR fragments are then run for fragment analysis, where each exon is represented by a defined length (which varies due to the stuffer sequence). This approach will reveal whether or not a given exon is present and allows the copy number of each exon to be calculated by comparing relative peak heights. MLPA can detect both deletions and duplications in patients and carriers. Small mutations within an exon can prevent binding of the probe to that exon, and as such, these mutations can present as single-exon deletions using the MLPA method.35 Therefore, a PCR using intronic primers is generally used to assess whether the exon is deleted or contains a small mutation. Furthermore, while MLPA detects which exons are involved in the deletion or duplication, it is not informative about the location of the intronic break points.

Alternatively, array comparative genome hybridisation (array CGH) uses probes covering dystrophin exons but also introns conjugated to a glass slide.36 Control and patient DNA is fragmented and labelled with separate fluorophores and hybridised to the probes. Like MLPA, this method will detect the relative abundance of each exon. However, because it also uses probes within introns, it allows pinpointing the location of the intronic breakpoint.

Multiplex PCR is not recommended for the genetic diagnosis of dystrophinopathy. However, it provides an alternative, cheaper approach to detecting deletions, and as such, is still used in numerous laboratories.37 38 This approach generally uses two primer sets to detect the presence or absence of exons 1, 3, 4, 6, 8, 12, 13, 17, 19, 43, 44, 45, 47, 48, 50, 51, 52 and 60. As such, it is able to pick up the majority of deletion mutations in patients with DMD and BMD. However, often the exact boundary of the mutation will be unknown. For example, when exon 45 is absent, this can be either an out-of-frame exon 45 deletion or an in-frame exon 45–46 deletion (because the presence of exon 46 is not assessed with this approach). This has ramifications for detecting whether a mutation is in-frame or out-of-frame, but also for assessing whether a patient is eligible for mutation-specific therapies like exon skipping. As such, the MLPA or array CGH are the recommended approaches to use first.33 In fact, historic cases identified with multiplex PCR in the past for whom the exact boundaries are not known should be reanalysed with MLPA because this can have implications for disease progression (in-frame vs out-of-frame mutations) and for eligibility for exon skipping therapy.

If no whole-exon deletions or duplications are found using MLPA, it is still possible that the patient has a small mutation in

**Figure 2** Schematic depiction of the effect of different types of Duchenne muscular dystrophy (DMD)-causing mutations on the dystrophin transcript. (A) Deletions of one or more exons can cause a shift of the open reading frame (in this example, a deletion of exon 45). (B) A duplication of one or more exons can cause a shift of the open reading frame (in this example, a duplication of exon 2). (C) There are several types of small mutations than can cause DMD. Nonsense mutations (top panel) introduce a stop codon prematurely (in this example, the nonsense mutation is located in exon 35). Small insertions or deletions (middle panel) can disrupt the open reading frame (in this example, a 1 bp insertion in exon 35). Finally, mutations affecting the splice sites (bottom panel) generally lead to the exclusion of the affected exon from the mRNA (in this example, exon 43). As such, a single-exon deletion that disrupts the open reading frame is generated on the mRNA level.
one of the 79 exons (20% of patients have small mutations). These can be identified using Sanger sequencing of individual exons, which is labour intensive and more expensive than MLPA or array CGH. Protocols for MLPA and Sanger sequencing for the DMD gene can be found on the Eurogentest website (http://www.eurogentest.org). It should be noted that not all laboratories will follow-up with exon analysis when MLPA (or multiplex PCR analysis) are negative. A recent survey was conducted among 41 individuals (primarily (paediatric) neurologists and clinical geneticists from Europe, Turkey and India) working with patients with DMD and attending a TREAT-NMD DMD masterclass (funded by an educational grant from PTC therapeutics) (survey results are being prepared for publication on http://www.treat-nmd.eu). This revealed that all respondents were aware of the need for a genetic diagnosis. However, when deletion and duplication tests are negative, >10% of responders would not perform confirmatory tests, while close to 90% would, using either exon sequencing or dystrophin analysis on a biopsy followed by exon sequencing. The main reason for not pursuing genetic analysis when deletion and duplication tests are negative was the costs involved in the Sanger sequencing or the lack of funding for shipping samples to laboratories able to conduct the small mutation analysis. It is possible that in the future MLPA, array CGH and Sanger sequencing will be replaced by next-generation sequencing techniques such as whole-exome sequencing for the genetic diagnosis of DMD. In fact, targeted sequencing of the DMD gene has already been reported using a library with probes covering the complete DMD gene to enrich for DMD and allowing a single-step diagnosis for deletions, duplications and small mutations in patients with DMD and BMD and carriers. However, currently it is more cost efficient to perform MLPA or array CGH followed by Sanger sequencing when no mutations are found. Obviously, there will be patients for whom no mutations are found. This can be because the mutation is located within an intron, as is the case for <0.5% of reported mutations, although these mutations may be under-represented because they are hard to detect. These mutations will be picked up with next-generation sequencing approaches, although without RNA analysis it may be challenging to assess whether intronic mutations

Table 2 Costs for genetic diagnosis of Duchenne muscular dystrophy in selected European countries

| Country      | Estimated cost (£) |
|--------------|--------------------|
| France       | 625                |
| Germany      | 3500               |
| Italy        | 700                |
| The Netherlands | 760            |
| Spain        | 1400               |
| Turkey       | 1600               |
| UK           | 1200               |

Estimated costs are based on personal communication with local clinical or molecular geneticists and neurologists Q1 2015. Costs include a full genetic diagnosis (so MLPA and if needed follow-up with Sanger sequencing).
lead to alterations on RNA level. Alternatively, the absence of a dystrophin mutation may indicate that the patient does not have DMD or BMD, but has a different form of muscular dystrophy.

Historically, the first step in diagnosing DMD was often protein analysis on a muscle biopsy by immunohistochemistry and/or western blot analysis. These techniques can reveal the size (western blot), location (immunohistochemistry) and abundance (both) of dystrophin, but in a diagnostic setting are most frequently used to show the presence or absence of dystrophin. Some laboratories perform a biopsy when MLPA analysis does not reveal deletions or duplications to confirm a dystrophinopathy before embarking on laborious small mutation analysis. However, because a genetic diagnosis is still required when absence of dystrophin is shown in a muscle biopsy, and because a muscle biopsy is an invasive procedure, the standards of care for DMD diagnosis suggest bypassing a muscle biopsy and using only genetic testing to diagnose DMD (figure 3). For most patients, muscle biopsies are never required. In exceptional cases, protein analysis can reveal whether dystrophin is absent in patients with a clear DMD phenotype but with no mutation in the DMD gene. If so, RNA can be isolated from the biopsied tissue and analysis may reveal an aberrant dystrophin mRNA transcript (eg, the inclusion of a cryptic exon due to a deep intronic mutation). Muscle biopsies may also be useful in patients who present with a discordant phenotype; for example, if a patient has an in-frame mutation but presents with DMD, a biopsy can reveal the absence of dystrophin. Vice versa, low levels of dystrophin detected in the muscle biopsy of a patient with an out-of-frame mutation presenting with a milder phenotype can explain this absence of dystrophin. Muscle biopsies may also be useful in patients who present with a dystrophinopathy before embarking on laborious small mutation analysis. However, because a genetic diagnosis is still required when absence of dystrophin is shown in a muscle biopsy, and because a muscle biopsy is an invasive procedure, the standards of care for DMD diagnosis suggest bypassing a muscle biopsy and using only genetic testing to diagnose DMD (figure 3). For most patients, muscle biopsies are never required. In exceptional cases, protein analysis can reveal whether dystrophin is absent in patients with a clear DMD phenotype but with no mutation in the DMD gene. If so, RNA can be isolated from the biopsied tissue and analysis may reveal an aberrant dystrophin mRNA transcript (eg, the inclusion of a cryptic exon due to a deep intronic mutation). Muscle biopsies may also be useful in patients who present with a discordant phenotype; for example, if a patient has an in-frame mutation but presents with DMD, a biopsy can reveal the absence of dystrophin. Vice versa, low levels of dystrophin detected in the muscle biopsy of a patient with an out-of-frame mutation presenting with a milder phenotype can explain this discrepancy. However, while dystrophin analysis may provide an explanation for why a disease course is slower or faster than expected, knowing this will not actually change the disease course. As such, this analysis is optional and not mandatory.

The standards of care for DMD were published in 2010 and timely genetic diagnosis are part of this. Early diagnosis for DMD is important because having genetic confirmation of the disease-causing mutation has important implications for the family. Once a mutation is identified, carrier analysis can be performed for the mother. With the exception of dystrophin, carrier analysis is performed on non-pregnant patients. For example, translocation (atuluren) has been approved in Europe for the treatment of DMD caused by nonsense mutations (in ambulatory patients aged 5 years or older). Given that DMD is a progressive disease and muscle loss is most likely irreversible, such disease-modifying treatments should be initiated as early as possible in eligible patients. Identification of the exact genetic mutation is also important for antiseizure-mediated exon skipping therapies, which are being tested in clinical trials and for which marketing authorisation applications are ongoing.

CONCLUDING REMARKS

DMD and BMD are both caused by mutations in the DMD gene. Having genetic confirmation of the mutation is important for patients because it has implications for disease prognosis, genetic counselling and evaluating each patient’s eligibility for emerging genetic therapies. Given that the progressive muscle wasting is irreversible, it is important that patients are identified as early as possible to consider all potentially effective treatments early in the disease course. On average, patients are being diagnosed at age 4.1 years in an expert centre compared with 4.5 years in 2000. Raising awareness with family physicians and other healthcare professionals who see young children could improve this further. Finally, given that new genetic therapies are emerging, one could consider offering neonatal screening for DMD.

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Competing interests AA-R declares being employed by LUMC, which has patents on exon skipping. As co-inventor of some of these patents, AA-R is entitled to a share of royalties. AAR is on the advisory board of ProQR and has been an ad hoc consultant for GLC consultancy, Global Guidepoint and PTC Therapeutics. Remuneration for these activities is paid to LUMC. KB declares being or being on the advisory board of Acceleron, AV Biopharma and Santhera Pharmaceuticals, being on the steering committee of BioMarin Nederland and being or having been a consultant for Dobiopharm, Genzyme, GSK, Prosensa Therapeutics/BioMarin Pharmaceutical, PTC Therapeutics, Lilly Pharmaceuticals, Pfizer, Summit Corporation, Insight Research Group, Galapagos SAGU, Shire Human Genetic Therapies Inc, Amsterdam Molecular Therapeutics, European Neuromuscular Centre, Bristol-Myers Squibb Company and Solid Ventures LLC.

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REFERENCES

1 Emery AE. The muscular dystrophies. Lancer 2002;359:687–95.
2 Flanagan KM. Duchenne and Becker muscular dystrophies. Neurol Clin 2014;32:671–88, viii.
3 Koenig M, Monaco AP, Kunkel LM. The complete sequence of dystrophin predicts a rod-shaped cytoskeletal protein. Cell 1988;53:219–28.
4 Aartsma-Rus A. Dystrophin analysis in clinical trials. J Neuromuscul Dis 2014;1:41–53.
5 Muntoni F, Torelli S, Ferlini A. Dystrophin and mutations: one gene, several proteins, multiple phenotypes. Lancet Neurol 2003;2:731–40.
6 Aartsma-Rus A, Van Deutekom JC, Fokkema IF, van Ommen GJ, den Dunnen JT. Entries in the Leiden Duchenne muscular dystrophy mutation database: an overview of mutation types and paradoxical cases that confirm the reading-frame rule. Muscle Nerve 2006;34:135–44.
7 Birkkannt DJ, Ararat E, Mhanna MJ. Cardiac phenotype determines survival in Duchenne muscular dystrophy. Pediatr Pulmonol 2015. http://dx.doi.org/10.1002/ppul.23215
8 Mendell JR, Shilling C, Leslie ND, Flanagan KM, al-Dahhak R, Gastler-Foster J, Kneile K, Dunn DM, Duval B, Aoyagi A, Hamil C, Mahmoud M, Roush K, Bird L, Rankin C, Lilly H, Street N, Chandrasekar R, Weiss RB. Evidence-based path to newborn screening for Duchenne muscular dystrophy. Ann Neurol 2012;71:304–13.
9 Moat SJ, Bradley DM, Salman R, Clarke A, Hartley L. Newborn bloodspot screening for Duchenne muscular dystrophy: 21 years experience in Wales (UK). Eur J Hum Genet 2010;21:134–53.
10 Heldereman-van den Enden AT, Madan K, Breuning MH, van der Hout AH, Bakker E, de Die-Smulders CE, Ginjaar HB. An urgent need for a change in policy revealed by a study on prenatal testing for Duchenne muscular dystrophy. Eur J Hum Genet 2013;21:21–6.
11 Abbas S, Tuffley-Giraud S, Bakker E, Ferlini A, Sejersen T, Mueller CR. Best practice guidelines on molecular diagnostics in Duchenne/Becker muscular dystrophies. Neuromuscul Disord 2010;20:422–7.
12 Bladen CL, Salgado D, Monges S, Foncuberta ME, Kekou K, Kosma K, Dawkins H, Lamont I, Roy AJ, Chamova T, Guergueltcheva V, Chan S, Kornkut L, Campbell C, et al. Aartsma-Rus A, et al. J Med Genet 2016;53:145–151. doi:10.1136/jmedgenet-2015-103387

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Aartsma-Rus A, et al. *J Med Genet* 2016;53:145–151. doi:10.1136/jmedgenet-2015-103387

Diagnostics

Dai Y, Wang J, Baricis N, Brabec P, Lahdetie J, Walter MC, Schreiber-Katz O, Karciagin V, Garamis V, Miswathanan V, Bayat F, Bucella F, Kufma E, Koeks Z, van den Bergen JC, Rodrigues M, Roxburgh R, Lusakowska A, Kostera-Pruszczyk A, Ignatkiewicz L, Eiwe VR, Emery AM, Peay HI, Bellard M, Kirschnere J, Flanigan KM, Straub V, Bushby K, Verschuuren J, Aartsma-Rus A, Beroud C, Lochmuller H. The TREAT-NMD DMD global database: analysis of more than 7,000 Duchenne muscular dystrophy mutations. *Hum Mutat* 2015;36:395–402.

Ankala A, Kohn JH, Hengke A, Meka A, Ephrem CL, Askreih SH, Bhide S, Hegre MR. Aberrant firing of replication origins potentially explains intrinsic non-coding rearrangements within genes, including the human DMD gene. *Genome Res* 2012;22:34–5.

Vulin A, Wein N, Strandjord DM, Johnson EK, Findlay AR, Maiti B, Howard MT, Kaminh Y, Taylor LE, Simmons TR, Ray WC, Montanaro F, Ervasti JM, Flanigan KM. The ZZ domain of dystrophin in DMD: making sense of missense mutations. *Hum Mutat* 2014;25:257–64.

Ray PN, Bellall D, Darte C, Logan C, Kean V, Thompson MW, Sylvester JE, Gorski JL, Kress W, Schara U. Xp21/A translocation: a rarely associated with Duchenne muscular dystrophy. *Nature* 1985;318:672–5.

Trippe H, Wieceorre K, Kottig W, Schara U. Xp21/A translocation: a rarely associated with Duchenne muscular dystrophy. *Muscle Nerve* 2015;45:333–5.

Ferreiro V, Giliberto F, Muniz GM, Francipane L, Marzese DM, Mampel A, Roque M, Barp A, Bello L, Politano L, Melacini P, Calore C, Polo A, Vianello S, Soraru G, Vo AH, McNally EM. Modi Neuron Neuropsych 2014;25:30–37.

Ginjaar IB, Kneppers AL, Meulen JD, Anderson LV, Bremmer-Bout M, Van Bergen JC, Ginjaar I, Niks E, Aartsma-Rus A, Verschuuren J. Prolonged ambulation following Duchenne and Becker muscular dystrophy. *PLoS ONE* 2015;10:e0141240.

Vo AH, McNally EM. Modifier genes and their effect on Duchenne muscular dystrophy. *Curr Opin Neurol* 2015;28:528–34.

Wein N, Vulin A, Falzarano MS, Szigayto CA, Maiti B, Findlay A, Keller NK, Uihlen M, Bakhvachavathala B, Messina S, Vita G, Passarelli C, Gualandi F, Wilton SD, Rodino-Klapac LR, Yang L, Dunn DM, Schoenberg DR, Weiss RB, Howard MT, Ferlini A, Flanigan KM. Translation from a DMD exon 5 IRES results in a functional dystrophin isoform that attenuates dystrophinopathy in humans and mice. *Nat Med* 2014;20:992–1000.

Winward AV, Mendell JR, Prior TW, Florence J, Burghele AS. Frame-shifted deletions of exons 3–7 revertant fibres in Duchenne muscular dystrophy; mechanisms of dystrophin production. *Am J Hum Genet* 1995;56:158–66.

Ginajba IK, Nepppe AS, Leuen JD, Anderson LV, Bremmer-Bout M, Van Duteke BM, Siegenewen J, Denuen NJ, Bakker E. Dystrophin nonsense mutations induce in-frame deletions of dystrophin resulting in variable exon 29 skipping and leads to variable phenotypes within one DMD family. *Eur J Hum Genet* 2000;8:793–6.

Bergen JC, Genaja JI, Nills E, Aartsma-Rus A, Verschueren J. Prolonged ambulation in Duchenne patients with a mutation amenable to exon 44 skipping. *J Neuromuscul Dis* 2015;21:1–9.

Panne M, Mazzone ES, Sormani MP, Messina S, Vita GL, Fanelli L, Baradelinni A, Tomente Y, D’Arco C, Langottti V, Viggiano E, D’Ambrosio P, Cavallaro F, Frosini S, Bello L, Bonfiglio S, Scalise R, De SR, Rolle E, Bianco F, Van der Haawe M, Magri F, Palermo C, Rossi F, Donati MA, Alfonsi C, Sacchini M, Arnoldi MT, Vita G, Compi GM, Ermani M, Calvo V, Angelini C, Hoffman EP, Pegoraro E. Genetic modifiers of Duchenne muscular dystrophy and diacylated dystrophin. *PLoS ONE* 2015;10:e0141240.

Vo AH, McNally EM. Modifier genes and their effect on Duchenne muscular dystrophy. *Curr Opin Neurol* 2015;28:528–34.

Genaja JI, Nills E, Aartsma-Rus A, Verschueren J. Prolonged ambulation in Duchenne patients with a mutation amenable to exon 44 skipping. *J Neuromuscul Dis* 2015;21:1–9.

Dai Y, Wang J, Baricis N, Brabec P, Lahdetie J, Walter MC, Schreiber-Katz O, Karciagin V, Garamis V, Miswathanan V, Bayat F, Bucella F, Kufma E, Koeks Z, van den Bergen JC, Rodrigues M, Roxburgh R, Lusakowska A, Kostera-Pruszczyk A, Ignatkiewicz L, Eiwe VR, Emery AM, Peay HI, Bellard M, Kirschnere J, Flanigan KM, Straub V, Bushby K, Verschuuren J, Aartsma-Rus A, Beroud C, Lochmuller H. The TREAT-NMD DMD global database: analysis of more than 7,000 Duchenne muscular dystrophy mutations. *Hum Mutat* 2015;36:395–402.

Ankala A, Kohn JH, Hengke A, Meka A, Ephrem CL, Askreih SH, Bhide S, Hegre MR. Aberrant firing of replication origins potentially explains intrinsic non-coding rearrangements within genes, including the human DMD gene. *Genome Res* 2012;22:34–5.

Vulin A, Wein N, Strandjord DM, Johnson EK, Findlay AR, Maiti B, Howard MT, Kaminh Y, Taylor LE, Simmons TR, Ray WC, Montanaro F, Ervasti JM, Flanigan KM. The ZZ domain of dystrophin in DMD: making sense of missense mutations. *Hum Mutat* 2014;25:257–64.

Ray PN, Bellall D, Darte C, Logan C, Kean V, Thompson MW, Sylvester JE, Gorski JL, Kress W, Schara U. Xp21/A translocation: a rarely associated with Duchenne muscular dystrophy. *Nature* 1985;318:672–5.

Trippe H, Wieceorre K, Kottig W, Schara U. Xp21/A translocation: a rarely associated with Duchenne muscular dystrophy. *Muscle Nerve* 2015;45:333–5.

Ferreiro V, Giliberto F, Muniz GM, Francipane L, Marzese DM, Mampel A, Roque M, Barp A, Bello L, Politano L, Melacini P, Calore C, Polo A, Vianello S, Soraru G, Vo AH, McNally EM. Modi Neuron Neuropsych 2014;25:30–37.

Ginjaar IB, Kneppers AL, Meulen JD, Anderson LV, Bremmer-Bout M, Van Bergen JC, Ginjaar I, Niks E, Aartsma-Rus A, Verschuuren J. Prolonged ambulation following Duchenne and Becker muscular dystrophy. *PLoS ONE* 2015;10:e0141240.