Multiple pathogenetic mechanisms in X linked dilated cardiomyopathy

N Cohen, F Muntoni

X linked dilated cardiomyopathy is a familial disease that is allelic to Duchenne and Becker muscular dystrophies and caused by mutations in the dystrophin gene. In several families with X linked dilated cardiomyopathy, the pattern of expression of dystrophin mutations in cardiac muscle differs from that in skeletal muscle. A number of these mutations affect transcription and splicing of the dystrophin gene in a tissue specific manner; others may affect regions of dystrophin that are presumed to have a more important role in cardiac than in skeletal muscle. These mutations are important because they highlight the fundamental differences in processing of the dystrophin gene between skeletal and cardiac tissues, as well as differences in the functional domains more relevant for one tissue or the other. This review focuses on the major mechanisms that have been proposed to explain this disorder.

Dilated cardiomyopathy (DCM) is a leading cause of cardiovascular morbidity and mortality. The DCMs form a heterogeneous group of disorders with different patterns of inheritance, including autosomal dominant (~23%), X linked (~5%), autosomal recessive, and mitochondrial transmission. About 30% of all DCMs are now considered to be inherited and the remaining 70% are sporadic. It is becoming increasingly clear, however, that a number of “sporadic” cases of DCM may be accompanied by de novo mutations in genes clearly associated with dominant disorders, complicating the issue of genetic counselling and prevalence of genetic disorders among patients with DCM. The best example is that of laminopathies, a significant contributor to the DCMs with conduction system disease, in which at least 50% of cases are the results of de novo mutations.

Five genes have so far been associated with X linked forms of cardiomyopathy and skeletal myopathy: tafazzin (also known as G4.5) gene,1 involved in Barth’s syndrome; the emerin gene,2 responsible for the X linked variant of Emery-Dreifuss muscular dystrophy; lysosome associated membrane protein 2 deficiency; involved in Danon’s disease; the XK membrane transport protein,3 responsible for McLeod’s syndrome;4 and the dystrophin gene,5 responsible for Duchenne (DMD) and Becker (BMD) muscular dystrophies and for X linked dilated cardiomyopathy (XLDC).6 Table 1 summarises the main features of these conditions.

Dystrophinopathy is the only condition that has been associated with exclusive cardiac involvement (XLDC). This form is secondary to mutations in the dystrophin gene, usually associated with DMD and BMD. DMD is the most common form of muscular dystrophy in childhood, occurring in one in 3500 male births. DMD is progressive, leading to loss of ambulation by 13 years. Although DMD can be inherited as an X linked condition, about one third of patients have de novo mutations. DMD is caused by a major reduction or absence in dystrophin expression at the sarcolemma.

Cardiac involvement in DMD develops insidiously during the first decade of life, at a time when skeletal muscle weakness is already significant.7 DCM is the most common type at all ages, occasionally following a mild hypertrophic phase. The clinical pathological aspects of cardiac involvement in DMD were recently reviewed by Finsterer and colleagues.8

Despite the high incidence of cardiac involvement, most patients with DMD remain asymptomatic. Respiratory failure caused by diaphragm muscle weakness is the most common cause of death in DMD, which generally occurs when patients reach their 20s.

BMD is a milder allelic form of DMD with an incidence of one in 14 000 live male births, but it is almost as common as DMD among the general population due to patients’ longer survival, usually well into adulthood. The clinical severity varies and, while patients at the severe end of the spectrum may be able to walk in their late teens, the majority remain ambulant for life.

Several studies have indicated that patients with BMD also have a high incidence of clinical cardiac involvement despite their milder skeletal muscle disease. In fact, some patients with BMD have initially presented with cardiomyopathy that became severe enough to require cardiac transplantation. The most common cause of death in BMD is heart failure. In a series of 68 patients with BMD studied from 1976 to 1993 whose diagnosis was confirmed by dystrophin testing, all had preclinical or clinical cardiac involvement by the age of 30.

Female carriers of DMD and BMD, an X linked recessive disorder, have a surprisingly high incidence of cardiac involvement that progresses with age and manifests primarily as cardiomyopathy. Cardiac muscle biopsies of DMD and

Abbreviations: BMD, Becker muscular dystrophy; DCM, dilated cardiomyopathy; DME, dystrophin muscle enhancer; DMD, Duchenne muscular dystrophy; XLDC, X linked dilated cardiomyopathy

See end of article for authors’ affiliations

Correspondence to: Professor F Muntoni, Dubowitz Neuromuscular Unit, Department of Paediatrics, Imperial College London, Hammersmith Hospital Campus, Du Cane Road, London W12 0NN, UK; f.muntoni@ic.ac.uk

Accepted 14 December 2003
BMD carries a mosaic pattern of absent (DMD) or reduced (BMD) dystrophin immunoreactivity mixed with cardiomyocytes with normal dystrophin expression. All of the conditions described above are associated with involvement of both skeletal and cardiac muscles. An intriguing example of cardiomyopathy, without any clinical sign of skeletal muscle weakness, is XLDC, an allelic condition to DMD and BMD. This article reviews the dystrophin gene mutations so far reported in patients with XLDC and summarises the proposed pathogenic mechanisms underlying the condition.

**XLDC**

XLDC was first described by Berko and Swift in 1987 in one large family with 11 affected young males and five manifesting carrier women, all of whom were mothers of affected sons. The condition was linked to the dystrophin gene in 1993 by Towbin and at the same time mutations in the dystrophin gene were identified by Muntoni and colleagues. No significant skeletal symptoms are present in this form. The clinical severity of the cardiac involvement in XLDC can be quite variable, ranging from an early onset and fatal cardiomyopathy to a milder form compatible with a better prognosis. The presentation, age at onset, and clinical course of the cardiomyopathy are variable. In the more common and severe form, patients typically develop congestive heart failure in their late teens or early 20s. There is rapid progression and a considerable risk of dying from congestive heart failure within a year from the initial presentation. Female carriers of XLDC, as in DMD and BMD, may also develop cardiomyopathy later on in life but the severity is usually milder, presumably due to mosaic expression of dystrophin in the heart. Increased serum concentrations of creatine kinase are often detected in patients and carriers.

Histological analysis of the skeletal muscle of patients with XLDC has shown a mild myopathy and variation in fibre size, with an increase in internal nuclei and interstitial connective tissue. In contrast, a severe dystrophic cardiomyopathy, characterised by notable fibrosis and variability in fibre size, is usually present in the cardiac muscle biopsies of patients with XLDC. A number of immunocytochemistry studies have shown that, although dystrophin expression is only mildly reduced in the patient’s skeletal muscle, it is absent in the cardiac muscle.

The first mutations identified in XLDC were clustered towards the 5’ end of the dystrophin gene and are the majority. However, mutations in other regions further downstream were subsequently identified. To date 16 different mutations in the dystrophin gene have been reported in patients with XLDC but the precise pathogenesis of this unique phenotype has remained unclear. This is partly due to the complexity of the dystrophin gene regulation within cardiac and skeletal muscles. In recent years, it has become clear that the frame shift hypothesis, based on the genomic structure of the gene after a deletion, cannot explain the cases of XLDC with mutations in the 5’ end of the gene. This theory, however, well accounts for the correlation between genotype and phenotype in most other cases of dystrophinopathy: in DMD the mutation typically induces the loss of the open reading frame but this is maintained in patients with BMD. Although the frame theory does not appear to provide an explanation for XLDC cases with mutations at the 5’ end, it accounts for a number of milder cases with in-frame deletions of the hot spot region.

On the basis of the region of the mutations, Tasaki and colleagues have recently classified patients with XLDC into three groups: those with mutations in the 5’ end of the gene (group I), or in the central hot spot region centred around exons 48–49 (group II), and those with mutations in regions other than those of group I and group II (group III). Since a number of mutations belonging to different “categories” affect splicing of the dystrophin gene (that is, the process in which the various exons are joined together and the introns removed from the mature messenger RNA), we propose a classification based on the likely mechanism of disease, which recognises two groups (table 2): group A includes mutations affecting transcription or splicing of the dystrophin gene resulting in a more severe cardiac involvement; group B includes mutations in which specific protein domains of dystrophin are affected.

**THE DYSTROPHIN GENE**

The dystrophin gene is the largest human gene known, spanning more than 3 Mb (that is, ~0.1% of the total human genome) or ~1.3% of the entire X chromosome. There are 79 exons and they account for ~0.6% of the gene, the residual part corresponding to intervening non-coding sequences (introns) and regulatory sequences.

Multiple transcripts are generated from the dystrophin gene, resulting in the formation of several isoforms. These derive from a combination of different tissue specific promoters and alternative splicing of the common dystrophin exons. There are at least seven different promoters. Three of these, specific for brain, muscle and Purkinje cerebellar cells, are arranged in series within the 5’ region of the gene and initiate transcription of full length messenger RNA (14 kb) each transcribing a novel exon 1 sequence. Figure 1 shows the promoter arrangement.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>X linked conditions responsible for cardiomyopathy and skeletal myopathy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disease</td>
<td>Gene</td>
</tr>
<tr>
<td>Barth’s syndrome (X linked recessive)</td>
<td>G4.4 (tafazzin)</td>
</tr>
<tr>
<td>DMD (X linked recessive)</td>
<td>Emerin</td>
</tr>
<tr>
<td>Danon’s disease (or XLCM-MR) (X linked recessive)</td>
<td>LAMP2</td>
</tr>
<tr>
<td>McLeod’s syndrome (X linked recessive)</td>
<td>XK</td>
</tr>
<tr>
<td>DMD (X linked recessive)</td>
<td>Dystrophin</td>
</tr>
<tr>
<td>BMD (X linked recessive)</td>
<td>Dystrophin</td>
</tr>
</tbody>
</table>

AV, atrioventricular; BMD, Becker muscular dystrophy; CK, creatine kinase; DCM, dilated cardiomyopathy; DMD, Duchenne muscular dystrophy; LAMP2, lysosome associated membrane protein 2; XLCM-MR, X linked vacuolar myopathy with cardiomyopathy and mental retardation.

www.heartjnl.com
XLDC and group A mutations

Most mutations causing XLDC are at the 5' end of the dystrophin gene. A deletion of the muscle promoter and part of the adjacent intron 1 has been described in a large Italian family and more recently in a Dutch family with XLDC. A point mutation abolishing the consensus 3' splice site of muscle exon 1 (that is, a mutation that would be expected to affect the splicing) has also been reported in two unrelated families. In another two unrelated Japanese families an insertion of a 5'-truncated form of human long interspersed nuclear element (LINE 1 or L1) transposable element in the 5'-untranslated region of the muscle exon 1 also resulted in XLDC. These three mutations display an identical transcription pattern in the skeletal muscle and, whenever studied, also in the cardiac muscle. A family with XLDC has been reported by Ortiz-Lopez in which a missense mutation in exon 9 of the dystrophin gene resulted in XLDC. In this family normal brain, muscle, and Purkinje isoform expression was detected in both cardiac and skeletal muscle.

Table 2  Literature review of dystrophin defect in XLDC patients

<table>
<thead>
<tr>
<th>Dystrophin gene mutation</th>
<th>CK</th>
<th>SK-MP</th>
<th>CA-MP</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Group A</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Muscle exon/intron 1 junction deletion</td>
<td>High</td>
<td>+</td>
<td>–</td>
<td>25</td>
</tr>
<tr>
<td>Muscle exon/intron 1 junction deletion</td>
<td>High</td>
<td>?</td>
<td>?</td>
<td>32</td>
</tr>
<tr>
<td>Muscle exon 1 3' splice site point mutation</td>
<td>Normal</td>
<td>+</td>
<td>–</td>
<td>28</td>
</tr>
<tr>
<td>Muscle exon 1 3' splice site point mutation</td>
<td>High</td>
<td>?</td>
<td>?</td>
<td>33</td>
</tr>
<tr>
<td>Splicing point mutation in intron 1</td>
<td>High</td>
<td>?</td>
<td>?</td>
<td>33</td>
</tr>
<tr>
<td>Insertion of L1 element in muscle exon 1</td>
<td>High</td>
<td>+</td>
<td>?</td>
<td>34</td>
</tr>
<tr>
<td>Insertion of L1 element in muscle exon 1</td>
<td>High</td>
<td>+</td>
<td>?</td>
<td>34</td>
</tr>
<tr>
<td>Exons 2–7 duplication</td>
<td>High</td>
<td>?</td>
<td>–</td>
<td>27</td>
</tr>
<tr>
<td>Exon 9 point mutation</td>
<td>High</td>
<td>+</td>
<td>–</td>
<td>38</td>
</tr>
<tr>
<td>Intron 11 deletion</td>
<td>High</td>
<td>+</td>
<td>–</td>
<td>26</td>
</tr>
<tr>
<td><strong>Group B</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exon 29 point mutation</td>
<td>High</td>
<td>+</td>
<td>+</td>
<td>39</td>
</tr>
<tr>
<td>Exon 35 missense mutation</td>
<td>High</td>
<td>?</td>
<td>?</td>
<td>33</td>
</tr>
<tr>
<td>Exons 45–48 deletion</td>
<td>Normal</td>
<td>?</td>
<td>?</td>
<td>37</td>
</tr>
<tr>
<td>Exons 45–55 deletion</td>
<td>High</td>
<td>?</td>
<td>?</td>
<td>31</td>
</tr>
<tr>
<td>Exon 48 deletion</td>
<td>High</td>
<td>?</td>
<td>?</td>
<td>36</td>
</tr>
<tr>
<td>Exon 48 deletion</td>
<td>Normal</td>
<td>+</td>
<td>–</td>
<td>37</td>
</tr>
<tr>
<td>Exons 48–49 deletion</td>
<td>High</td>
<td>?</td>
<td>?</td>
<td>35</td>
</tr>
<tr>
<td>Exons 48–51 deletion</td>
<td>High</td>
<td>?</td>
<td>+</td>
<td>37</td>
</tr>
<tr>
<td>Exons 48–53 deletion</td>
<td>Normal</td>
<td>?</td>
<td>+</td>
<td>37</td>
</tr>
<tr>
<td>Exons 49–51 deletion</td>
<td>Normal</td>
<td>+</td>
<td>+</td>
<td>35</td>
</tr>
</tbody>
</table>

+, present; –, absent; ?, not known; CA-MP, dystrophin protein in cardiac muscle; L1, long interspersed nuclear element; SK-MP, dystrophin protein in skeletal muscle.

Figure 1  Physical map of the dystrophin gene. Green boxes indicate exons, blue boxes indicate exons specific for each promoter (B1, first brain exon; M1, first muscle exon; P1, first Purkinje exon), and red boxes indicate promoters (BP, brain promoter; MP, muscle promoter; PP, Purkinje promoter; S1, Schwann cell promoter). Not drawn to scale.
skeletal muscle tissues. However, there were no studies on the effect of this mutation on gene transcription. The observation that dystrophin protein was absent in the heart but reduced in skeletal muscle suggests that the mutation may result in tissue specific splicing of the dystrophin gene in this family.

Our group has also reported an unusual mutation involving deletion of dystrophin intron 11.26 In this family, a deletion of intron 11 has brought two naturally occurring cryptic splice sites (hidden splice sites) that are usually ignored by the splicing machinery close together, thus activating them. This results in the incorporation of part of dystrophin intron 11 as a novel exon into dystrophin mRNA transcript, abolishing the normally spliced transcript in the cardiac muscle. As this interrupts the reading frame, it results in the absence of dystrophin in the cardiac muscle. However, the novel exon coexists with normally spliced exon 11 to exon 12 transcript in the skeletal muscle allowing a significant residual expression of the protein (30% compared with normal control).26 Transcriptional analysis of the skeletal and cardiac muscle of this family showed an essentially normal pattern of isoform expression in the heart and skeletal muscle.26 We have found that normally spliced transcript is the only dystrophin transcript detected in patient lymphocytes. This finding not only indicates a tissue specific splicing pattern for the dystrophin gene in this family but also highlights the unpredictable nature of lymphocytes in identifying splicing mutations in patients with dystrophinopathies.

**XLDC and group B mutations**

Franz and colleagues reported on a family with XLDC25 in which a nonsense mutation in exon 29 caused skipping of this exon in both cardiac and skeletal muscle.27 Skipping is the mechanism by which an exon that would normally be incorporated in the mature transcript is missed out. In this patient, dystrophin is severely reduced in both skeletal and cardiac muscle.28 Recently, Feng and colleagues reported a missense mutation of exon 35 causing a substitution of a highly conserved asparagine by lysine in a patient with XLDC.29

Mutant dystrophin transcripts have not been well characterised in patients with mutations involving a central hot spot region but all the deletions so far identified in these patients are in frame and typically associated with milder BMD. Two patients affected by XLDC have been reported to carry in-frame deletions of exons 49–51 in one and 48–49 in the other. In the former patient almost normal concentrations of truncated protein were detected in the heart.30 Clusters of deletions of exons 45–48, exon 48, exons 48–51, and exon 48–53 have also been reported.31 In these patients reduced but preserved dystrophin expression was detected in the cardiac muscle.32 Recently, in-frame deletion of exons 45–55 has been described in a patient with XLDC.33

In contrast to group A patients, in whom dystrophin is totally absent within the heart, in group B patients the dystrophin concentration is reduced and no major difference in promoter usage or splicing between skeletal and cardiac muscle has been reported. Thus, in group B the mutations cause a qualitative rather than a quantitative defect in perturbing the expression of dystrophin. This suggests that a mechanism(s) other than aberrant splicing is involved in this group of patients with XLDC, possibly affecting the structure of dystrophin.

**PROPOSED PATHOGENIC MECHANISMS**

The finding of different mutations, dystrophin mRNA processing, and protein expression in these cases suggests that multiple pathogenic mechanisms account for all XLDC.

**Mutations affecting the splicing of the dystrophin gene (group A)**

Various regions of the dystrophin gene seem to be involved in this group of mutations.

Interestingly, up to 43% of dystrophin gene mutations that cause XLDC specifically affect expression of the muscle isoform. Transcription studies in all XLDC patients with mutations at the 5’ end of the dystrophin gene have detected up regulation of the brain and Purkinje isoforms of dystrophin in skeletal muscle but not in cardiac muscle.34 35 36 37 This up regulation compensates for the lack of the muscle isoform. This observation further underlines the important role of both the brain and Purkinje isoforms in maintaining stability of the cytoskeletal organisation in skeletal muscle, thus offering protection from progressive degeneration of the muscles. Up regulation of the brain and Purkinje isoforms has also been reported in the skeletal muscle of a patient with BMD with typical skeletal muscle involvement and a deletion affecting the first muscle exon 1 and the first intron intron 1.38 Nakamura and colleagues proposed that the brain and Purkinje promoters may respond to reduced dystrophin expression in these patients. In addition, up regulation of the two promoters has also been detected in four patients with DMD with deletion not involving the 5’ end of the dystrophin gene.39 These observations suggest that the observed up regulation is not limited to patients with XLDC or mutations at the 5’ end of the gene. However, up regulation of the brain and Purkinje promoters would only provide an advantage to patients in whom there is no mutation in downstream exons but the defect is limited to the muscle promoter or first muscle exon.

Activation of the brain and Purkinje promoters may depend on the presence of DNA regulatory elements that may become functionally relevant under specific circumstances. In this respect, conserved DNA motifs present in the regulatory regions of several muscle specific genes have been described in the brain and Purkinje promoters.40 41 Sequence analysis of the dystrophin brain and Purkinje regions identified domains that were highly homologous to CAAG boxes and CARG boxes plus E boxes, respectively; these motifs are binding sites for the myogenic determining gene (Myod, myf5, myogenin, MRF4) and serum response factor families of transcription factors.42–45 Recently, the role of a dystrophin muscle enhancer 1 (DME1) in up regulation of the brain and Purkinje isoforms in muscle cell has been shown.32 This enhancer is preserved in two of the patients with mutation involving muscle exon/intron junction deletion.46 More recently, Bastianutto and colleagues identified DME2, which contained cardiac specific transcriptional elements. This enhancer apparently has a role in dystrophin gene regulation at later stages of cardiac muscle development. The presence of two other possible enhancer elements, DME3 and DME4, has also been predicted.47 Figure 2 shows the arrangement of DMEs.

In the family described by Bies and colleagues in which exons 2–7 were duplicated, the same mutant dystrophin transcript was spliced in both cardiac and skeletal muscle tissues but probably with different efficiency.48 The mutation may have caused a cardiac sensitive change in dystrophin function only. Since the dystrophin actin binding domain is essential for dystrophin function, the disruption of this interaction may lead to an exclusive impairment of cardiac muscle function. However, part of the difference was apparently also due to an alteration of dystrophin abundance in the heart, resulting from differences in transcription between cardiac and skeletal muscle. Thus, we hypothesise that a combination of disruption within the dystrophin cardiac domain and tissue specific splicing of dystrophin has a significant role in pathogenesis of XLDC in this family.
The mutation reported by Ferlini and colleagues involving an intronic 11 deletion is clearly a splicing mutation, as incorporation of the novel pathogenic exon is regulated in a tissue specific manner. One of the interesting features of this mutation is that all the exons are intact and thus an entirely intronic deletion is involved in XLDC. This observation suggests the presence of cardiac specific regulatory sequences within the dystrophin intron. Intronic enhancers have been shown to be involved in regulation of alternatively spliced exons in cell specific, differentiation stage, and tissue specific manners. These sequences presumably work by recruiting splicing factors to the 5′ splice site. Many examples of inhibitory cis-acting elements (genes that are located on one chromosome and act on or cooperate with neighbouring genes on the same chromosome) have also been described, showing that splice sites can be blocked by secondary structure or by specific or general factors binding to regulatory intronic elements near the repressed sites.

Although some of the mutations reported in this group definitively affect splicing and transcription, in others there was not sufficient tissue to perform parallel studies in cardiac and skeletal muscle. An example of these cases is found in the family described by Ortiz-Lopez and colleagues with a missense mutation of exon 9. The authors suggest that the mutation affecting the conserved first hinge region of the protein is likely to result in loss of membrane integrity and eventual loss of function. This was probably due to the continual stress placed on the beating pump, therefore favouring the hypothesis of an important functional role for the mutated domain. However, no detailed studies on the effect of this mutation on gene transcription in this region could be carried out. It is important to note that the A to G substitution occurred at nucleotide +4 of exon 9, located within one exonic consensus splicing sequence. Since the 5′ and 3′ exonic extremities are involved in splicing accuracy and efficacy, it is possible that this mutation affects splicing mechanisms. In addition, the rare physiological occurrence of exon 9 skipping suggests that transcription studies in this patient may be very important for ruling out a splicing pathology.

**Mutations affecting the dystrophin domain are particularly important for its cardiac function and interaction (group B)**

The great majority of patients belonging to this group carry mutations in the spectrin-like domain of dystrophin. There is no clear explanation as to why cardiac muscle is preferentially affected in them. Three possibilities can be hypothesised: firstly, trans-acting factors (protein products produced by genes on one chromosome and cooperating with or acting on genes elsewhere) in the cardiac muscle of patients with XLDC may cause misregulation of dystrophin splicing, ultimately affecting dystrophin expression in the heart; secondly, intronic cis-acting dystrophin specific sequence(s) may be lost; and thirdly, some specific domains that have a specific or prevalent cardiac function may be lost.

Although the first hypothesis appears to be very unlikely, we will analyse here the available evidence in favour of or against it. In keeping with the idea that alternative splicing may have a role in these patients, Sironi and colleagues have recently found at least seven alternative splicing events that are involved in the dystrophin gene encompassing exons 44–58 in healthy skeletal muscle. They also noticed that deletions involving the same exons can determine diverse splicing behaviours in different patients or even different tissues of the same person. A study of alternative splicing events in large number of patients with BMD, carried out by the same group, suggests that altered splicing patterns in these patients do not directly derive from the gene defect but may derive from trans- rather than cis-acting factors. Furthermore, recent observations in up regulation of a trans-acting factor in DMD skeletal muscle has led to the proposal of the role of such factors in splicing regulation. Unfortunately, transcription has not been systematically studied in patients with XLDC with group B mutations. However, we analysed the transcription of dystrophin in the cardiac muscle of a patient with a deletion in this region and found a normal pattern of splicing. Therefore, neither the available transcription data nor the available protein expression data support the hypothesis that differences in splicing between cardiac and skeletal muscle have a significant role in these patients.

Regarding the second hypothesis, that the loss of intronic regions with potential regulatory functions may have a role in patients with these deletions, since the incidence of cardiac involvement is high in patients with BMD with deletions involving exon 49 but not those with deletions of only exon 48, it has been proposed that intron 48 may contain sequences important to the function of dystrophin in cardiac muscle. Owing to the very large size of the intron, this hypothesis has been difficult to prove.

In keeping with the “epitope” theory (the third hypothesis), the central hot spot region of the gene contains hinge regions (fig 3) that may be important for a specific function or interaction of dystrophin in the heart. Recently it has been shown that the dystrophin domain encoded by exons 45–48 is required for localisation of neuronal nitric oxide synthase to the sarclemma. The authors suggested that this may relate to conformational changes induced by mutations in this region of dystrophin or to a yet unidentified protein interaction in this region. While neuronal nitric oxide synthase is not expressed in cardiac muscle, this study provided evidence that this region of dystrophin is involved in the interaction with dystrophin associated proteins. It can therefore be hypothesised that additional proteins relevant for heart function may interact with this domain of dystrophin.

In the family reported on by Franz and colleagues, a nonsense mutation in exon 29 results in exon skipping and a significant reduction of dystrophin in both cardiac and skeletal muscle. The detection of aberrant splicing in both tissues makes it difficult to understand the different mechanisms involved in protein expression of these tissues. On the basis of the dystrophin interaction with actin and the sarcosomal dystrophin associated glycoprotein complex, Franz and colleagues have hypothesised that conformational change in the mid-region of the dystrophin may profoundly affect this interaction and cause membrane instability. The rod-like domain of dystrophin is formed by 24 tandemly repeated segments, each consisting of three α helices connected by turns. Exon 29 encodes most of two of the helices and one of the turns. The deletion of exon 29

---

**Figure 2** A model of dystrophin gene with muscle enhancer elements. DME, dystrophin muscle enhancer.
utrophin, which is very similar in overall structure to dystrophins, including chicken and dog fish, and in region involved in this mutation is highly conserved in conserved asparagine by lysine in a patient with XLDC. The mutation of exon 35 causing substitution of a highly dystrophin deficient mice are more susceptible to enterovirus change in the mid-rod region.\textsuperscript{39} There is also increasing evidence for secondary changes in cardiomyopathy, and in particular dystrophin may provide a final role in function and interaction of dystrophin in the heart.

Recently, Feng and colleagues\textsuperscript{13} reported a missense mutation of exon 35 causing substitution of a highly conserved asparagine by lysine in a patient with XLDC. The region involved in this mutation is highly conserved in dystrophins, including chicken and dog fish, and in utrophin, which is very similar in overall structure to dystrophin.\textsuperscript{33} On the basis of the observed conservation of the mutated amino acid, the second hypothesis may be suggested. However, since there has been no report of a transcriptional study for this patient it is difficult to rule out other mechanisms.

In addition to a primary role of dystrophin in XLDC, in recent years a potential link between this protein and environmental conditions has also been identified. Dystrophin is proteolytically cleaved by coxsackieviral protease 2A, functionally impaired, and morphologically disrupted in cultured cardiomyocytes as well as in the intact mouse heart infected with coxsackievirus B3, and this has been proposed to contribute to the pathogenesis of inflammatory DCM.\textsuperscript{69} In addition it has been reported that the hearts of dystrophin deficient mice are more susceptible to enterovirus induced cardiomyopathy.\textsuperscript{68}

There is also increasing evidence for secondary changes in dystrophin in heat failure. Recently, Vatta and colleagues\textsuperscript{2A} have shown that decreased contractile function is associated with selective disruption of the amino terminus of dystrophin in patients with end stage cardiomyopathy (idiopathic dilated or ischaemic). The authors hypothesised that an unidentified protein may interact with and stabilise the amino terminus of dystrophin. It is this interaction that is disrupted in end stage heart failure. This disruption was reversible in some patients after left ventricular assistance device support. These observations support the hypothesis that changes in cytoskeletal proteins and in particular dystrophin may provide a final common pathway for contractile dysfunction in heart failure and that these changes may well be reversible by reduction of mechanical stress.\textsuperscript{69}

**SUMMARY**

We have summarised the mutations identified so far in patients with XLDC. A number of these mutations affect transcription, splicing, or both in the dystrophin gene in a tissue specific manner. Others are more likely to be involved in destabilising regions of dystrophin and have a more important role in cardiac than in skeletal muscle. Overall, these mutations are important, as they highlight the fundamental differences in the processing of the dystrophin gene between skeletal and cardiac tissues, as well as the functional domains more critical for one tissue or the other. Recent evidence also suggests that dystrophin may have a significant role in acquired infective or ischaemic cardiomyopathies. Further studies are necessary for understanding the underlying pathogenesis of DCM.

**ACKNOWLEDGEMENTS**

This work was supported by the British Heart Foundation. The authors thank Afshin Cohen for reading the manuscript.

**Authors’ affiliations**

N Cohen, F Muntoni, Dubowitz Neuromuscular Unit, Department of Paediatrics, Imperial College London, Hammersmith Hospital Campus, London, UK

**REFERENCES**

Pathogenesis of X-linked dilated cardiomyopathy


