Electrophysiological abnormalities in induced pluripotent stem cell-derived cardiomyocytes generated from Duchenne muscular dystrophy patients

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Abstract
Duchenne muscular dystrophy (DMD) is an X-linked progressive muscle degenerative disease, caused by mutations in the dystrophin gene and resulting in death because of respiratory or cardiac failure. To investigate the cardiac cellular manifestation of DMD, we generated induced pluripotent stem cells (iPSCs) and iPSC-derived cardiomyocytes (iPSC-CMs) from two DMD patients: a male and female manifesting heterozygous carrier. Dystrophin mRNA and protein expression were analysed by qRT-PCR, RNAseq, Western blot and immunofluorescence staining. For comprehensive electrophysiological analysis, current and voltage clamp were used to record transmembrane action potentials and ion currents, respectively. Microelectrode array was used to record extracellular electrograms. X-inactive specific transcript (XIST) and dystrophin expression analyses revealed that female iPSCs underwent X chromosome reactivation (XCR) or erosion of X chromosome inactivation, which was maintained in female iPSC-CMs displaying mixed X chromosome expression of wild type (WT) and mutated alleles. Both DMD female and male iPSC-CMs presented low spontaneous firing rate, arrhythmias and prolonged action...
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1 | INTRODUCTION

DMD caused by mutations in the dystrophin gene, is an X-linked muscle degenerative disease present in 1.7-2.1 of 10 000 male births.1-3 Homozygous symptomatic females are rare and heterozygous females are mostly unaffected carriers, yet ~20% are symptomatic manifesting carriers with later and milder disease progression than male patients.4-6 In DMD patients dystrophin mutations lead to destabilization of the dystrophin glycoprotein complex (DGC) and sarcolemma instability resulting in functional deficits.7 Dystrophin also plays a role in anchoring dystrophin-associated proteins to the sarcolemma, including nitric oxide synthase (nNOS).8 Absence of functional dystrophin ultimately leads to degeneration and death of skeletal and cardiac muscle, being replaced by fibrous tissue.9 DMD symptoms begin at early childhood, mainly including muscle fatigue. With age, the disease progresses and wheelchair assistance is needed in early teen years. Eventually, respiratory function is compromised and requiring mechanical support. Cardiac dysfunction usually manifests during late teenage or early twenties, but ultimately dilated cardiomyopathy (DCM) and congestive heart failure develop with ventricular and supraventricular arrhythmias. Finally, patients progress to death because of respiratory or cardiac failure.10

In female embryos, one of the X-chromosomes is randomly inactivated leading to mosaicism of maternal and paternal X-linked alleles explaining why heterozygous females are usually less affected or asymptomatic. iPSCs enable generation of human disease models to study underlying pathomechanisms in a cell type specific level. Previous studies investigating the effect of reprogramming on X chromosome inactivation status suggest three possible outcomes: (1) X chromosome inactivation (XCI) status is maintained throughout the reprogramming process; (2) inactivated X chromosome is reactivated; (3) inactivated X chromosome is eroded.11-18

To investigate the electrophysiological abnormalities caused by dystrophin mutations in cardiomyocytes, we generated iPSCs from two DMD patients: a male and a female heterozygous manifesting carrier. X inactivation status and dystrophin gene expression were analysed in iPSCs and iPSC-CMs generated from healthy controls and patients of both sexes with regard to genetic, molecular and functional abnormalities. In support of the hypothesis that dystrophin-mutated iPSC-CMs from the mutant male and the heterozygous female carrier exhibit key features of DMD, mutated cardiomyocytes displayed molecular, genetic and electrophysiological abnormalities, including arrhythmias.

2 | METHODS

See Supporting Information for detailed Methods.

Dermal biopsies were obtained from a 50-year-old DMD female manifesting patient carrying a deletion of exons 8-12 (ex.8_12del) and a 32-year-old DMD male patient carrying a substitution of cytosine to thymine (c.5899C>T) constituting a premature stop codon. The donors signed a consent form according to approval #7603-09-SMC by the Helsinki Committee for Experiments on Human Subjects at Sheba Medical Center, Ramat Gan, Israel. See Supporting Information for Methods including, patients’ clinical history, iPSCs generation and characterization,19 karyotype analysis, teratoma formation, genotyping and pluripotency evaluation by immunofluorescence staining and flow cytometry. Differentiation into cardiomyocytes was performed according to the directed differentiation by modulating Wnt/β-catenin signalling as previously described.20 Transmission electron microscopy was used to evaluate ultrastructural features of iPSC-CMs. See Supporting Information for protein and RNA expression analyses.

2.1 | Electrophysiological experiments

Action potentials were recorded from small cardiomyocyte clusters and single cells in whole-cell configuration. The pacemaker current \( I_f \) was recorded from single cardiomyocytes (enzymatically dissociated) in the presence of 500 μM BaCl₂ to block \( I_{K1} \). To record \( I_f \), the membrane was clamped at 15 seconds intervals, from a holding potential of −40 mV to −120 mV in 10 mV steps for 2 seconds pulse durations.21 To record L-Type Ca\(^{2+}\) current \( (I_{Ca,L}) \), the membrane was clamped at 200 ms depolarizing steps, from a holding potential of −70 mV ranging from −40 to +40 mV after a 20 ms −40 mV pulse for inactivation of Na\(^+\) currents.22 Axopatch 200B, Digidata 1322 or
The male DMD patient suffered from muscular dystrophy from early childhood and was diagnosed with DCM at the age of 17. At age 25, he was hospitalized with respiratory insufficiency primarily because of combined respiratory and heart failure, and required tra-
cheostomy and prolonged ventilation. At that time, his LVEF was 15% and the ECG showed sinus rhythm with narrow QRS and QE pattern in L1, AVL and V2-3. At age 30, he was respirator-dependent but his heart failure was reasonably controlled. The echo-Doppler showed LVEDD 58 mm, LVEF 20%, moderate mitral insufficiency and normal right ventricular function. At age 33, a routine Holter-ECG showed for the first time episodes of non-sustained ventricular tachycardia of 3-12 beat at a rate of up to 200/minute. Two years later, because of repeated episodes of ventricular flutter deteriorating into ventricular fibrillation, in order to save his life an ICD was implanted.

The female patient, a manifesting carrier, presented at age 42 with proximal muscle weakness and creatine kinase elevation. Her son diagnosed with DMD expired at age 16. A year later, her echo-Doppler showed a biventricular dysfunction with left ventricular dimension of 65 mm, left ventricular ejection fraction of 30% and moderate to severe mitral insufficiency. Her ECG showed low voltage with left axis deviation and narrow QRS, she suffered from paroxysms of atrial fibrillation with rapid ventricular response and non-sustained ventricular tachycardia. An ICD was implanted, but despite heart failure therapy, her condition continued to deteriorate. Cardiorespiratory exercise test showed VO₂max of 6 ml/kg/min (33% of normal value) resulting from primarily cardiovascular limitation, but the patient declined heart transplantation. At age 49, she was in NYHA IV. AV nodal ablation and upgrading to CRTD was required because of atrial fibrillation with rapid ventricular response. Echo-Doppler showed severe biventricular dysfunction with LVEF of 20% and severe tricuspid regurgitation. The muscle weakness progressed to complete inability to walk. The patient expired at age 51 because of end-stage heart failure associated with renal insufficiency.

3.2 Molecular characterization of DMD iPSCs and iPSC-CMs

iPSCs were generated from dermal fibroblasts. For iPSCs characterization, FACS and immunostaining validated pluripotent markers expression (Figures S1 and S2). Teratoma assay demonstrated iPSCs ability to differentiate into the three germ layers (Figure S1). Karyotype was preserved throughout the reprogramming process, and Sanger sequencing confirmed the mutations in the patients iPSCs (Figure S1). For control, iPSCs and iPSC-CMs were generated from four different healthy female and male donors with varying genetic backgrounds, thus markedly strengthening the comparisons between the mutated and control populations.

iPSCs differentiation resulted in cardiomyocytes and myofibrillarogenesis (Figure 1A) depicted by troponin I expression (at 30 days) and spontaneous contractions. Ultrastructural analyses did not reveal any differences between the experimental groups, including changes in sarcomere length (Figure S4). Using dystrophin exon-specific qRT-PCR, control and DMD male iPSC-CMs produced robust induction of full-length 427 kD muscle isoform of dystrophin mRNA, while total dystrophin transcript measured by a common C-terminal exon only increased modestly overtime post-differentiation. Utrophin mRNA expression was also induced during differentiation but was suppressed during late iPSC-CMs development. Muscle-specific γ-sarcoglycan was also expressed following differentiation in control and DMD male iPSC-CMs. By Western blotting (WB), full-length 427 kD dystrophin was detected in control and DMD female iPSC-CMs, but undetectable in DMD male iPSC-CMs, confirming a protein null allele in this patient (Figure 1C). All clones manifested slow skeletal Tnl (ssTnl), the neonatal isoform of troponin I in iPSC-CMs indicating induction of sarcomerogenesis. To test assembly of dystrophin with the DGC, membrane preparations were prepared from iPSC-CMs, solubilized and passed over wheat germ agglutinin affinity columns, which demonstrated that dystrophin co-purified with glycoprotein fractions similar to adult heart tissue from normal and Becker muscular dystrophy patients with a truncated dystrophin allele (Figure 1D).
is a long non-coding RNA on the X chromosome that acts as a major effector of the X inactivation process.\textsuperscript{17,28} Expression analysis demonstrated the absence of XIST RNA in iPSCs, indicating that XCR or erosion of XCI occurred during reprogramming of DMD and control female dermal fibroblasts. In agreement with previous reports,\textsuperscript{24,25,29} XIST expression patterns were maintained during differentiation of iPSCs into iPSC-CMs (Figure 1G). As XIST does not fully correlate with XCI in iPSCs,\textsuperscript{17,30} thus preventing distinction between XCR and erosion of XCI, X chromosome status could not be determined definitively. To investigate the expression pattern of dystrophin in iPSC-CMs, we used next generation sequencing (NGS) which demonstrated (Figure 1E,F) mixed dystrophin expression in the female iPSC-CM population. Specifically, dystrophin transcripts expressed as mRNA fragments per kilobase of exon per million reads mapped (FPKM) values were normalized by library size and corrected by enrichment factor. Black arrow indicates zero expression of the mutated allele in control iPSC-CM samples. Control n = 5; DMD female, n = 5. Two-way ANOVA followed by Holm-Sidak post-hoc analysis. ***P < 0.001. (F) Sashimi plot of the WT allele RNA in control iPSC-CMs (top) and both WT and mutated alleles RNA in DMD female iPSC-CMs (bottom). (G) Expression of XIST RNA in female compared to male human fibroblasts (HF), iPSC and iPSC-CM as determined by qRT-PCR. Expression of XIST was normalized to β-actin and relative expression levels are shown as 2$^{-\Delta\Delta CT}$ compared to the respective male control
3.4 | Electrophysiological abnormalities in female and male DMD iPSC-CMs

Common pathophysiological phenomena in DMD patients are supraventricular and ventricular arrhythmias, the latter being the second most common cause of death in these patients. Accordingly, we investigated action potential properties and specific ionic currents in control and DMD iPSC-CMs generated from multiple independent differentiations. A prominent electrophysiological abnormality in DMD iPSC-CMs was slower spontaneous firing rates compared to control cardiomyocytes; female—21.6 bpm, male—17.6 bpm, control—47.4 bpm (Figures 2, 3A). To investigate the mechanism underlying this finding, we compared the properties of I\textsubscript{f} (a key pacemaker current) in control versus DMD iPSC-CMs. As illustrated by representative current traces (Figure 4A) and I-V relations (Figure 4B), while DMD female iPSC-CMs did not differ from control, I\textsubscript{f} density in DMD male iPSC-CMs was smaller (P < 0.001). No differences were observed in current activation relations of the three groups (Figure 4C), implying similar channel properties. To decipher the mechanisms underlying the differences in I\textsubscript{f} I-V relations among the groups, we measured the expression of the hyperpolarization-activated cyclic nucleotide-gated channel—isoforms 2 and 4 (HCN2 and HCN4) by means of qRT-PCR analysis. These experiments demonstrated increased expression of HCN2 and HCN4 in DMD female iPSC-CMs compared to control and DMD male iPSC-CMs (Figure 4D).

3.4.1 | Arrhythmias in DMD iPSC-CMs

In addition to the slow firing rate, DMD female and male iPSC-CMs displayed voltage oscillations, which resemble delayed afterdepolarizations (DADs) and oscillatory prepotentials (OPPs) during action potential recordings. As demonstrated in Figure 2, DMD female and male iPSC-CMs presented oscillatory depolarization in 52% and 17% of spontaneous recordings, respectively (Figure 2), while control iPSC-CMs had no arrhythmias.

3.4.2 | Action potential parameters

As seen in Figure 3, action potential parameters of DMD female and male iPSC-CMs differed from control. Mean action potential durations at 20/50/90% repolarization (APD\textsubscript{20}/APD\textsubscript{50}/APD\textsubscript{90}) were longer than control; further, the mean APD\textsubscript{20}/APD\textsubscript{50}/APD\textsubscript{90} were longer in DMD male than in female iPSC-CMs (Figure 3F-H). Importantly, markedly prolonged APD\textsubscript{90} (>400 ms) was observed in 29% and 100% of DMD female and male iPSC-CMs, respectively. Finally, mean maximal diastolic potential was more negative in DMD male iPSC-CMs (~69.6 mV) than control (~62.6 mV) (Figure 3B).

To neutralize the effect of slow firing rate in DMD cardiomyocytes on APD, we measured action potential parameters in paced single cell cardiomyocytes. In agreement with the results from spontaneously firing cardiomyocytes, APD\textsubscript{20}/APD\textsubscript{50}/APD\textsubscript{90} were prolonged in DMD female and male iPSC-CMs (Figure 5A, 5-E), while other parameters were comparable to control (Figure 5B). To identify the specific action potential phase responsible for APD prolongation, we calculated the ratio between APD at different stages of repolarization and APD\textsubscript{90}; the APD contributing mostly to prolongation was APD\textsubscript{20}, implying involvement of early repolarization phase (ie APD\textsubscript{20}; Figure 5B).

3.4.3 | Increased I\textsubscript{Ca,L} density in DMD iPSC-CMs

To determine the mechanism underlying APD\textsubscript{20} prolongation, we measured I\textsubscript{Ca,L} the major depolarizing current in early repolarization. As illustrated by the representative current traces (Figure 6A) and mean I-V relations (Figure 6B), I\textsubscript{Ca,L} density was larger in both DMD female and male than in control iPSC-CMs (female—at voltage steps from 0 mV to −20 mV and male—at 0 mV). No differences were observed in the current activation curve among the three groups (Figure 6C). In agreement with larger I\textsubscript{Ca,L} density, qRT-PCR analysis demonstrated increased expression of the Ca\textsuperscript{2+}-voltage-gated channel subunit alpha1 C (CACNA1C) in DMD female and male compared to control iPSC-CMs (Figure 6D).

3.4.4 | Differences in the BRV between male and female patients

BRV representing the dynamic non-linear features of automaticity was analysed in spontaneous action potential recordings (1000-1800 sec) (Figure 7A). Whereas DMD female iPSC-CMs displayed more dispersed inter-beat interval (IBI) scatter plot and histogram than control (Figure 7B-D), DMD male iPSC-CMs IBI scatter plot and histogram were similar to control. Representative Poincaré plots, a quantitative measure of the evolution in time of the BRV, demonstrate the firing pattern abnormalities in DMD female and male iPSC-CMs compared to the typical ellipsoid-shaped control Poincaré plot (Figure 7F-I). Furthermore, coefficient of variation (CV) and SD2, referring to long term IBI variability, were increased in DMD female iPSC-CMs compared to control (Figure 7J-L). DMD male iPSC-CMs CV and SD2 were similar to control (Figure 7J-L). Overall, DMD female iPSC-CMs displayed increased BRV measures including IBI CV and SD2 Poincaré plot index compared to control iPSC-CMs. Importantly, similar results were obtained at the network level, from analysis of extracellular electrogrograms recorded by means of micro-electrode array (MEA; Figure S5).

4 | DISCUSSION

In support of the hypothesis that dystrophin-mutated iPSC-CMs exhibit key features of DMD disease, our main findings were: (1) WB analysis showed expression of full length 427 kD dystrophin in control and DMD female iPSC-CMs, but not in DMD male iPSC-CMs, confirming that the male patient carries a null allele; (2) XIST and dystrophin expression analyses showed that iPSC generation from female dermal cells is associated with transcriptional reactivation of the inactivated X chromosome either through XCR or erosion
of XCI, which is maintained during iPSC-CM differentiation; (3) DMD female iPSC-CM population expresses both WT and mutated dystrophin alleles; (4) DMD female and male iPSC-CMs display arrhythmias and APD prolongation.

4.1 | Unique genetic features of DMD female iPSC-CMs population exhibiting mosaic expression of DMD alleles

DMD female carriers are usually unaffected by dystrophin mutations because of skewed XCI preferably silencing the mutated allele. However, in this study we generated iPSC-CMs from a heterozygous female manifesting carrier who presented severe adult onset DCM and myopathy. Importantly, this woman carried a highly malignant out-of-frame deletion in the 5′ terminal of the DMD gene which caused a severe disease that resulted in teenage death of her son. Genetic analysis demonstrates the DMD female iPSC-CM population mixed expression of WT dystrophin allele alongside the mutated allele (Figure 1). During reprogramming, X chromosome status is destined to be one of the following three options: (1) conserved inactivation pattern of the originating somatic cell (XaXi); (2) reactivation of the inactive X chromosome (XaXa); (3) erosion of XCI (XaXe).

Considering XIST expression does not sufficiently indicate X chromosome status, and based on the NGS dystrophin expression analyses,
a likely possibility is that a mixed population was established during reprogramming and each differentiation. In this scenario, DMD female iPSC-CMs can be divided to two sub-populations: (1) cells expressing the WT allele, and (2) cells expressing the mutated allele. Nonsense mediated decay of mRNA containing premature stop codon, reduces the amount of detectable mutated mRNA in the samples, thus hampering exact quantification of the ratio between these two sub-populations. Nevertheless, our genetic analysis faithfully represents the two dystrophin alleles expression ratio in the iPSC-CM population used for the functional experiments. Furthermore, these novel results display an abnormally increased expression of WT allele in the iPSC-CM population, implying the first sub-population (expressing the WT allele) also carries an abnormality compared to control iPSC-CMs.

4.2 | Electrophysiological abnormalities in DMD iPSC-CMs

DMD iPSC-CMs exhibit a variety of electrophysiological abnormalities divided into four categories: (1) decreased automaticity and altered $I_f$ density; (2) arrhythmias including DADs and OPPs; (3) prolonged APD and enhanced $I_{Ca,L}$ density; (4) increased BRV in DMD female iPSC-CMs.

4.2.1 | Decreased automaticity and altered $I_f$ density

Contrary to DMD patients who usually manifest increased high heart rate, DMD iPSC-CMs display low spontaneous firing rate...
compared to control. However, in DMD patients, increased heart rate is attributed to elevated sympathetic tone; hence, as this masking effect is absent in DMD iPSC-CMs, the in vitro study exposes a fundamental pathophysiology of the pacemaker function — attenuated automaticity. To elucidate the mechanism underlying this reduced automaticity in the male and female cardiomyocytes, we recorded $I_f$, a major pacemaker current. In agreement with the reduced automaticity in DMD male cardiomyocytes, $I_f$ density was smaller than control, albeit similar mRNA expression levels of HCNs. In DMD female iPSC-CMs $I_f$ density was similar to control, while mRNA expression levels of HCNs were significantly increased. Overall, both DMD female and male iPSC-CMs demonstrate reduced $I_f$ density relatively to their respective mRNA expression levels of HCNs. The precise mechanism involved in altered HCN function is yet to be determined. No differences were observed in current activation between DMD and control iPSC-CMs.

### 4.2.2 Arrhythmias including DADs and OPPs

In contrast to regular firing pattern in control iPSC-CMs, 52% and 17% of DMD female and male cardiomyocytes exhibited arrhythmias including DADs and OPPs, respectively. The common cause for DADs is intracellular Ca$^{2+}$ overload leading to Na$^+$/Ca$^{2+}$ exchanger activation resulting in transient inward depolarizing current ($I_h$) and DADs initiation. The proposed mechanisms underlying Ca$^{2+}$ overload in DMD cardiomyocytes are: (1) damaged sarcolemma enables penetration of extracellular Ca$^{2+}$; (2) Inflammatory mediators increase the expression of INOS which binds to and destabilizes the...
Ryanodine receptors (RyRs) located on the sarcoplasmic reticulum (SR), resulting in SR Ca\(^{2+}\) leakage into the cytosol.\(^{38}\) Notably, leaky RyRs, which can lead to Ca\(^{2+}\) overload, were reported in mdx mice.\(^{39}\) Furthermore, \(I_{Ca,L}\) was increased in DMD iPSC-CMs compared to control iPSC-CMs (Fig. 6). The phenomenon of DADs and OPPs in DMD iPSC-CMs is in agreement with increased Ca\(^{2+}\) influx and Ca\(^{2+}\) overload, which can contribute to cardiac arrhythmias in DMD patients.\(^{40}\)

4.2.3 | Prolonged APD and enhanced \(I_{Ca,L}\) density

Importantly, the prolonged APD in female and male DMD cardiomyocytes, potentially associated with fatal arrhythmias,\(^{41}\) may account for clinical phenotypes observed in DMD patients, such as long QT interval (LQT).\(^{42}\) The ratio between the changes in APD\(_{20}\) and APD\(_{90}\) strongly implies that most of the prolongation stems from early repolarization phase. As \(I_{Ca,L}\) is the major depolarizing current...
during early repolarization,33 we measured its density which was higher in both DMD female and male iPSC-CMs. These results are consistent with previous findings of enhanced $I_{Ca,L}$ in adult cardiomyocytes of mdx mice.43 Furthermore, CACNA1C expression was increased in both DMD female and male iPSC-CMs compared to control.

4.2.4 Increased BRV in DMD female iPSC-CMs

DMD female (but not male) iPSC-CMs displayed increased BRV measures compared to control cardiomyocytes. While further investigation is required to determine the contribution of the mixed dystrophin expression to this finding, it is likely that heterogeneity of the cardiomyocytes expressing either WT or mutated dystrophin alleles, contributes to increased BRV indices in action potential recordings from iPSC-CM clusters.

4.3 Advantages of DMD iPSC-CMs over the mdx mouse model

The common murine model for DMD research is the mdx mouse which carries a point mutation in exon 23 of the dystrophin gene.43 However, the similarity of mdx hearts to human DMD hearts is debated; while mice initially develop hypertrophic cardiomyopathy and only at a later stage contractile dysfunction and systolic heart failure, human cardiomyopathy is characterized primarily by ventricular dilatation and reduced systolic function.44 The use of patients' iPSC-CMs has been previously demonstrated as valid cellular in vitro model for various genetic heart diseases including DMD,45–48 as well for LQT syndrome,49 and PRKAG2 myopathy.50 Furthermore, iPSC-CMs are an accessible human cellular model which also enables exclusion of in vivo masking factors such as the influence of the autonomous nervous system.

4.4 Summary and conclusions

In conclusion, this study demonstrates that loss of dystrophin in patients’ iPSC-CMs is sufficient to cause electrophysiological abnormalities including low spontaneous firing rate, arrhythmogenic firing patterns, abnormal action potential parameters, increased BRV and altered pacemaker currents density. Importantly, this is the first study demonstrating the impact of mixed dystrophin expression pattern on abnormal characteristics and functional abnormalities of DMD iPSC-CMs generated from a DMD female manifesting carrier. These results may suggest greater predisposition to arrhythmias in early stages of DMD cardiomyopathy in females. Further studies are therefore required to characterize the natural history and define the arrhythmic risk in female patients with DMD mutation. This study also has implications for genetic therapies aimed at restoring dystrophin expression by gene editing or gene therapy; if genetic correction is incomplete, a subset of dystrophin deficient myocytes may increase electrophysiological heterogeneity which could be a substrate for arrhythmogenesis.

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CONFLICT OF INTEREST

The authors have nothing to declare.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.