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Effects of cardioactive drugs on human induced pluripotent stem cell derived long QT syndrome cardiomyocytes

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Abstract

Human induced pluripotent stem cells (hiPSC) have enabled a major step forward in pathophysiologic studies of inherited diseases and may also prove to be valuable in in vitro drug testing. Long QT syndrome (LQTS), characterized by prolonged cardiac repolarization and risk of sudden death, may be inherited or result from adverse drug effects. Using a microelectrode array platform, we investigated the effects of six different drugs on the electrophysiological characteristics of human embryonic stem cell-derived cardiomyocytes as well as hiPSC-derived cardiomyocytes from control subjects and from patients with type 1 (LQT1) and type 2 (LQT2) of LQTS. At baseline the repolarization time was significantly longer in LQTS cells compared to controls. Isoprenaline increased the beating rate of all cell lines by 10–73 % but did not show any arrhythmic effects in any cell type. Different QT-interval prolonging drugs caused prolongation of cardiac repolarization by 3–13 % (cisapride), 10–20 % (erythromycin), 8–23 % (sotalol), 16–42 % (quinidine) and 12–27 % (E-4031), but we did not find any systematic differences in sensitivity between the control, LQT1 and LQT2 cell lines. Sotalol, quinidine and E-4031 also caused arrhythmic beats and beating arrests in some cases. In summary, the drug effects on these patient-specific cardiomyocytes appear to recapitulate clinical observations and provide further evidence that these cells can be applied for in vitro drug testing to probe their vulnerability to arrhythmia.

Keywords: Induced pluripotent stem cell, Patient-specific, Long QT syndrome, Cardiomyocytes, Multielectrode array, Cardioactive drug, Arrhythmia

Background

Inherited long QT syndrome (LQTS) is a potentially severe arrhythmic disease that affects the electrical repolarization of the myocardium and manifests as an abnormally long QT interval on electrocardiogram (ECG) recordings. LQTS may result in polymorphic ventricular tachycardias known as torsades de pointes (TdP), which can ultimately lead to life-threatening ventricular fibrillation and sudden cardiac death (Schwartz et al. 2013).

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Inherited forms of LQTS typically result from mutations in cardiac ion channel coding genes. In the case of LQT type 1 (LQT1), the mutations are present in the KCNQ1 gene which encodes the α -subunit of the slow component of the delayed rectifier potassium current (K_v 7.1) (Wang et al. 1996), whereas LQT2 is caused by mutations of the KCNH2 gene which encodes the α -subunit of the rapid component of the delayed rectifier potassium current (K_v ,11.1) (Sanguinetti et al. 1995). Four founder mutations in these two genes have been found to explain the high (0.4 %) prevalence of LQTS in the Finnish population (Marjamaa et al. 2009). The most prevalent of these founder mutations is the C-terminal KCNQ1 G589D missense mutation, resulting in a dysfunctional assembly domain and preventing the formation of fully functional potassium channel tetramers (Piippo et al.

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2001). Another prevalent but apparently milder founder mutation, the *KCNH2* R176W missense mutation, resides in the N-terminus of the channel and accelerates the deactivation of the I_{Kr} tail current, thus leading to reduced potassium efflux (Fodstad et al. 2006). Research on the exact effects of these mutations on the functional properties of cardiomyocytes has been hampered by the lack of suitable techniques that can adequately recapitulate the full spectrum of the disease in vitro.

Human induced pluripotent stem cell (hiPSC) technology has recently made it possible to produce human cell lines with characteristics similar to those of embryonic stem cells from somatic cells of various origins (Takahashi et al. 2007; Yu et al. 2007). To date, multiple disease-specific hiPSC lines have already been established (Wu and Hochedlinger 2011). Several studies have focused on LQTS using the hiPSC technology (Bellin et al. 2013; Itzhaki et al. 2011; Kiviaho et al. 2015; Lahti et al. 2012; Matsa et al. 2011, 2014; Moretti et al. 2010). These studies suggest that hiPSC technology may be useful for modeling LQTS in vitro. Until now, large-scale drug effects have been studied with human embryonic stem cell-derived cardiomyocytes (hESC-CMs) and wildtype hiPSC-CMs (Braam et al. 2010; Caspi et al. 2009; Harris et al. 2013; Mehta et al. 2013; Navarrete et al. 2013; Yokoo et al. 2009). In this study, we investigated the effect of six clinically used drugs in control hESC- and wildtype hiPSC-CMs and in hiPSC-derived LQT-CMs. Additional value will be gained by studying drug responses at clinically relevant concentrations (Schulz and Schmoldt 2003), since many of the mutation carriers can have a latent, subclinical form of LQTS that may be exacerbated upon pharmacological challenge (Roden 2004).

We created hiPSC lines from the dermal fibroblasts of patients with *KCNQ1* G589D and *KCNH2* R176W mutations. We investigated how these patient-specific hiPSCs

respond to pharmacological challenges in comparison with cardiomyocytes derived from control hESCs and hiPSCs.

Methods

Patient-specific pluripotent stem cell induction

The LQT1- and 2-specific hiPSCs were induced as previously described (Takahashi et al. 2007) from fibroblasts of patients carrying the *KCNQ1* (G589D) or the *HERG* (R176W) mutation (Fodstad et al. 2004) (see Additional file 1: Materials and methods). The study has been approved by Pirkanmaa Hospital District ethical committee (R08070). Details of the hiPSC lines are listed in Table 1. The subjects volunteered for the study gave their consent for skin biopsy to be taken.

Stem cell culture

The control pluripotent stem cell lines used in this study were H7 (hESC, WiCell), UTA.00112.hFF [fetal wild-type from ATCC (WT)-hiPSC], and UTA.01006.WT (WThiPSC derived from a healthy adult). For LQT1, 4 patientspecific hiPSC lines were used: UTA.00208.LQT1, UTA.00211.LQT1, UTA.00313.LQT1 and UTA.00303. LQT1. For LQT2 2 patient-specific hiPSC lines were used: UTA.00514.LQT2 and UTA.00525.LQT2. The pluripotent stem cells were cultured on a mouse embryonic fibroblast (MEF, Millipore) feeder cell layer in KSR medium consisting of KnockOut Dulbecco's Modified Eagle's Medium (KO-DMEM, Invitrogen) supplemented with 20 % KnockOut serum replacement (KO-SR, Invitrogen), 1 % non-essential amino acids (NEAA, Lonza), 1 % Glutamax (Invitrogen), 50 U/mL penicillin/streptomycin (Lonza), and recombinant human basic fibroblast growth factor (R&D Systems). The cells were passaged once a week by treating the pluripotent stem cell colonies with collagenase IV (Gibco) and seeding them onto a fresh MEF feeder layer.

Table 1 Details of the control and patient-specific pluripotent stem cell lines used for drug testing

Cell line	Line type	Mutation	Disease	Affected current	Source	Purpose hESC control	
H7 (hESC)	hESC	Wild type	None	None	ICM		
UTA.00112.hFF (WTa)	hiPSC	Wild type	None	None	hFF	Control	
UTA.01006.WT (WTb)	hiPSC	Wild type	None	None	hADF	Adult control	
UTA.00208.LQT1 ^{0,*,s}	hiPSC	G589D	LQT1	I _{Ks}	hADF	Symptomatic, patient-specific	
UTA.00211.LQT1 ^{0,*,s}	hiPSC	G589D	LQT1	I _{Ks}	hADF	Symptomatic, patient-specific	
UTA.00303.LQT1 ^{ψ,*,a}	hiPSC	G589D	LQT1	l _{Ks}	hADF	Asymptomatic mutation carrier-specific	
UTA.00313.LQT1 ^{ψ,*,a}	hiPSC	G589D	LQT1	I _{Ks}	hADF	Asymptomatic mutation carrier-specific	
UTA.00514.LQT2 ^{‡,a}	hiPSC	R176W	LQT2	l _{Kr}	hADF	Asymptomatic mutation carrier-specific	
UTA.00525.LQT2 ^{‡,a}	hiPSC	R176W	LQT2	I _{Kr}	hADF	Asymptomatic mutation carrier-specific	

ICM inner cell mass of blastocyst, hFF human foreskin fibroblast, hADF human adult dermal fibroblast, hESC human embryonic stem cell, hiPSC human induced pluripotent stem cell

* Siblings $^{\ddagger, \, \theta, \, \psi}$ clonal lines from same individual s symptomatic a asymptomatic

Cardiomyocyte differentiation

Cardiomyocytes were differentiated from hESCs as well as from the control and LQT1 and -2-specific hiPSCs using the mouse visceral endoderm-like (END-2) cell coculture method, as described elsewhere (Mummery et al. 2003). Briefly, mitomycin C (Sigma) -inactivated END-2 cells were trypsinized and plated onto 12-well plates in 0 % KO-SR hES medium. The pluripotent stem cells were plated on top of the END-2 cell monolayer the following day. The 0 % KO-SR hES medium was refreshed on days 5, 8 and 12. At day 14, the medium was changed to 10 % KO-SR hES medium, and it was refreshed three times a week thereafter.

Pharmacological testing

Drugs

Cisapride monohydrate, quinidine and (\pm) -sotalol hydrochloride were obtained from Sigma-Aldrich and dissolved in dimethyl sulfoxide (Sigma-Aldrich) at 10 mM. E-4031 (Alomone Labs) and erythromycin (Abboticin i.v., Amdipharm) were dissolved in sterile H₂O at 1 and 50 mg/mL, respectively. Isoprenaline (Isuprel i.v., Hospira) was supplied in ready-to-use ampoules. The drug concentrations were chosen based on their therapeutic blood serum concentration range (Schulz and Schmoldt 2003). The following concentrations of cisapride monohydrate, quinidine, (\pm) -sotalol hydrochloride, and erythromycin were tested: half the concentration of the lower limit for the therapeutic blood serum concentration, the lower limit of the therapeutic blood serum concentration range, the average therapeutic serum concentration, the upper limit of the therapeutic range, and twice the upper limit of the therapeutic concentration (Schulz and Schmoldt 2003). The isoprenaline concentrations were chosen according to previously published values and multiples thereof (Pekkanen-Mattila et al. 2009). The drugs were serially diluted in embryoid body (EB)medium with 5 % fetal bovine serum (FBS, see below). The test concentrations for each drug are listed in Table 2.

Table 2 Drugs and their concentrations used with cardio-myocytes derived from human embryonic and inducedpluripotent stem cells

Drug	Concentrations
Isoprenaline	nM: 81, 402
Cisapride monohydrate	nM: 40, 83, 120, 165, 330
Erythromycin	μM: 1.5, 3, 5.5, 8, 16
Sotalol	μM: 0.8, 1.6, 5.7, 9.7, 19.4
Quinidine	μM: 1.5, 3, 9, 15, 30
E-4031	nM: 10, 100, 300, 500, 700
Quinidine E-4031	μM: 0.0, 1.0, 3.7, 5.7, 13.1 μM: 1.5, 3, 9, 15, 30 nM: 10, 100, 300, 500, 700

Multielectrode array recordings

The hESC-CMs and the control, LQT1-, and LQT2specific hiPSC-CM aggregates were mechanically excised from the cultures and plated onto fetal bovine serum (PAA) and 0.1 % gelatin- (Sigma-Aldrich) coated 6-well-MEAs (6wellMEA200/30iR-TI-mr, MultiChannel Systems MCS GmbH). The FPs were recorded with a USB-MEA1060 amplifier, and the temperature was kept at +37 °C using a TC02 heating element (both from Multi Channel Systems MCS GmbH). The MEAs were covered with gas-permeable MEA membranes (ALA MEA-SHEET, ALA Scientific) during the recordings. A wash-in time of 2 min was allowed for the drugs before recordings. The beating areas were cultured in EB medium consisting of KO-DMEM (Invitrogen) supplemented with 20 % FBS (Invitrogen), 1 % NEAA (Lonza), 1 % Glutamax (Invitrogen), and 50 U/mL penicillin/streptomycin (Lonza). The drug tests were performed in 5 % FBS containing EB medium; the medium was changed at least an hour prior to the baseline recordings, and the cells were allowed to stabilize in the incubator (+37 $^{\circ}$ C, 5 % CO₂). The field potential signals were recorded with MC Rack v.4.0.0 software (Multi Channel Systems MCS GmbH).

Data and statistical analysis

The field potential durations (FPDs) from the recorded files were measured offline using our in-house developed CardioMDA software (Pradhapan et al. 2013) and AxoScope10 (Molecular Devices). The baseline data are presented as mean \pm standard deviation (SD). Linear mixed-effect models with either Bazett's corrected field potential duration (cFPD) or beating rate (BR) as a dependent variable were fitted using the function lme in R (Software environment for statistical computing and graphics, version 2.13.0, The R Foundation for Statistical Computing). The factorial variable for the concentration level was used as an independent variable. The lowest concentration level was used as a reference group. Random intercept for different aggregates was used together with independent random errors.

Results

LQT patient characteristics

Skin biopsies were obtained from two siblings, a 28-year old female asymptomatic LQT1 patient (QTc interval, 428 ms) and a 41-year old female symptomatic LQT1 patient (QTc interval, 456 ms), both with the *KCNQ1* G589D mutation. In addition, skin biopsies were obtained from a 61-year old male asymptomatic LQT2 patient with the *KCNH2* R176W mutation (QTc interval, 437 ms). The symptomatic LQT1 patient had experienced seizures before beta-blocker medication, while the asymptomatic sibling has never had any cardiac

symptoms. The asymptomatic LQT2 individual has only experienced occasional palpitations, which although are not specific symptoms of LQTS.

Molecular characteristics of cardiomyocytes

Reverse-transcription PCR and immunofluorescence microscopy showed that the control and LQT-hiPSCs expressed the pluripotency markers NANOG, OCT3/4, SOX2, SSEA4, TRA-1-60, and TRA-1-81 (Additional file 1: Fig. S1A, Table S1). Furthermore, the hiPSC colony morphology resembled those of hESCs, the hiPSCs differentiated into cardiomyocytes at efficiencies previously reported for hESCs upon END-2 co-culture (data not shown) (Pekkanen-Mattila et al. 2009), showed nuclear localization of the cardiac-specific transcription factor NKX2.5 which is also expressed in hESC-CMs (Asp et al. 2010), and exhibited a distinct striated pattern in cardiac troponin T staining (Additional file 1: Fig. S1B). The electrophysiological properties and drug responses were recorded with microelectrode arrays (MEAs).

Effects of cardioactive drugs to the cFPD of hiPSC-CMs

We compared the drug responses of the control cardiomyocytes derived from hESC line H7 (hereafter called hESC) and two wild-type hiPSC lines UTA.00112.hFF (hereafter called WTa) and UTA.01006.WT (hereafter called WTb) to the hiPSC-derived LQT-CMs (for the cell line properties, see Table 1). The results of the clonal cell lines derived from the same individual were pooled together (UTA.00208.LQT1 and UTA.00211.LQT1 [=LQT1A], UTA.00303.LQT1 and UTA.00313.LQT1 [=LQT1B], UTA.00514.LQT2 and UTA.00525.LQT2 [=LQT2]). LQT1A were derived from a symptomatic patient, LQT1B from an asymptomatic mutation carrier and LQT2 from asymptomatic mutation carrier.

Baseline characteristics

The baseline beating rates (BR) showed large differences between cell populations, ranging from 59 to 105 beats per minute (Fig. 1a). However, these BRs did not differ significantly among cardiomyocytes differentiated from hESCs, control-, or LQT-hiPSCs. In contrast, the beating rate-corrected field potential duration (cFPD, Bazett's) showed significant differences between control- and LQT-CMs (Fig. 1b). At baseline level, the LQT-CMs showed significantly longer cFPD (586–600 ms) compared to control-CMs (340–400 ms) (p < 0.001).

Isoprenaline effects

To test whether the differentiated cardiomyocytes had proper β -adrenergic responses, we subjected them to an isoprenaline challenge. This treatment had chronotropic effect on cardiomyocytes and increased the BR at the

concentration of 81 and 402 nM (Fig. 2). Isoprenaline did not induce any arrhythmic beats in any of the CMs, including those derived from LQT1- and LQT2-hiPSC lines.

Effects of non-cardiac drugs cisapride and erythromycin

Cisapride increased the repolarization time in all CMs studied, although this effect did not reach statistical significance in the LQT2-CMs, apparently due to large experimental variation (Fig. 3a).

Erythromycin likewise induced a significant average increase of cFPD in all cell lines studied, in a concentration range of 1.5–16 μ M (Fig. 3b). No arrhythmic beats or beating arrests were observed with cisapride or with erythromycin.

Effects of the anti-arrhythmic drugs sotalol and quinidine

Upon sotalol (0.8–19.4 μ M) challenge, the maximal cFPD prolongation ranged from of 8 % (WTb-CMs) to 23 % (LQT1A-CMs) (Fig. 4a). Quinidine also caused a concentration-dependent (1.5–30 μ M) average cFPD prolongation in all the cardiomyocytes (Fig. 4b). Of all clinically used drugs tested (cisapride, erythromycin, sotalol, quinidine), quinidine showed the largest cFPD-increasing effect (31–42 %) seen in LQT1-CMs.

The effect of the hERG channel blocker E-4031

E-4031 also caused concentration-dependent cFPD prolongation in all cardiomyocyte types (Fig. 4c). It is of note that in hESC-CMs all but one of the cardiac aggregates became arrhythmic or stopped beating at concentrations of 500–700 nM. Thus, error bars indicating standard deviation could not be calculated for these experimental points. The cFPD prolongation in the different cell lines reached statistical significance at E-4031 concentration of 100 nM indicating that the sensitivities of the different cell lines to E-4031 were very similar (Fig. 4c).

In summary, Fig. 5 illustrates the representative traces of cisapride, erythromycin, sotalol, quinidine and E-4031 in control- and LQT-CMs.

The proarrhythmic effects of sotalol, quinidine and E-4031

Sotalol, quinidine and the hERG blocker E-4031 showed proarrhythmic effects on certain individual cell lines (Table 3). With sotalol, only spontaneous afterdepolarization (AD)-like waveform abnormalities (but not beating arrests) were observed (Table 3). Sotalol caused AD-like waveforms ranging from concentrations 5.7 μ M (average therapeutic serum concentration) to 19.4 μ M (twice the upper therapeutical serum concentration). None of the LQT1B-CMs showed any arrhythmic behavior upon sotalol challenge.

Quinidine showed also beating arrests in addition to AD-like waveform abnormalities (Table 3). The AD-like

waveforms were observed at the therapeutical serum concentration range (3–15 μ M) in hESC-, LQT1A- and LQT2-CMs but not in WT- or LQT1B-CMs (Table 3). On the other hand, the beating arrests occurred ranging from 9 μ M (average therapeutic serum concentration) to 30 μ M (twice the upper therapeutical serum concentration) (Table 3).

The compound E-4031 also caused AD-like waveform abnormalities and beating arrests (Table 3). E-4031 caused AD-like waveform abnormalities in cardiomyocytes differentiated from all of the lines except WTb-CMs. Furthermore, beating arrests were seen in all cardiomyocyte types except LQT1A-CMs. Among the LQTs, the highest frequency for AD-like waveform abnormalities was found in LQT1A-CMs (50 % of cardiomyocyte aggregates). On the other hand, the highest frequency for beating arrests were seen in LQT2-CMs (31 % of cardiomyocyte aggregates).

Discussion

The present study further demonstrates the usefulness of the hiPSC-derived cardiomyocytes to serve as a model





for long QT syndrome, showing that the QT intervalprolonging effects of clinically used drugs are replicated in this in vitro system, using concentrations that are within or close to their therapeutic range (Schulz and Schmoldt 2003). We showed that LQT-CMs corrected field potential duration is significantly longer than in control-CMs at baseline conditions, which suggests that long QT syndrome is recapitulated in these cells. This study also reproduces in cell clusters the earlier observation with single cells by patch clamp with prolonged APDs in LQT-CMs compared to controls (Kiviaho et al. 2015; Lahti et al. 2012). In the current study LQT- and control CMs were further exposed to various cardioactive drugs to investigate their effects.

The cardiomyocytes beating rates were increased by isoprenaline, which is a β -adrenergic agonist that affects calcium ($I_{Ca,L}$) channel activation via β_1 receptors. This is in line with the previous studies showing positive chronotropic effect of isoprenaline on both hESC- and hiPSC-CMs (Moretti et al. 2010; Pekkanen-Mattila et al. 2009; Yokoo et al. 2009). β -adrenergic stimulation with isoprenaline did not have any arrhythmic effects on the LQT1 lines although LQT1 patients often experience symptoms at increased heart rates.

Cisapride is a serotonin 5-HT4 receptor agonist that has been reported to block the hERG potassium channel (Mohammad et al. 1997). It is known to induce TdP in patients (Darpö 2001). Caspi et al. (2009) reported significant (>50 nM) dose-dependent prolongation of cFPD in hESC-CMs (Braam et al. 2010; Caspi et al. 2009).

We obtained similar results as Caspi et al. (2009) as the hESC-CMs used in our study showed dose-dependent and significant cFPD prolongation starting at 83 nM. In contrast, Braam et al. (2010) reported no significant cisapride-induced cFPD prolongation in hESC-CMs. However, they judged the therapeutic concentration to be 2.6–4.9 nM (Redfern et al. 2003), whereas we chose the therapeutic range as 83-165 nM (Schulz and Schmoldt 2003), which may explain the fact that they were not able to detect significant cisapride-induced cFPD prolongation. Mehta et al. (2013) have also investigated the effect of cisapride to the cFPD of 150-170 days old EB-based hiPSC-derived CMs. Likewise, they also found concentration-dependent cFPD prolongation (Mehta et al. 2013). Mehta et al. (2013) did not observe any arrhythmias with cisapride, which is in line with our study here.

Erythromycin is a macrolide antibiotic that prolongs the QT interval in patients (Shaffer et al. 2002). The therapeutic plasma concentration of erythromycin has been reported to be $3-8 \mu$ M (Schulz and Schmoldt 2003). In our study, erythromycin prolonged significantly cFPD in hESC-CMs already at the first test concentration (1.5 μ M). Our data may suggest a slightly higher sensitivity of LQT2-CMs compared to LQT1-CMs, but high experimental variation precludes firm conclusions. Erythromycin did not induce any arrhythmic effects in the cardiomyocytes.

Sotalol is a β -adrenergic receptor antagonist and a Vaughan–Williams class III anti-arrhythmic agent that potently blocks the potassium ion channels in



cardiomyocytes (Edvardsson et al. 1980). In hESC-CMs, Braam et al. (2010) reported cFPD prolongations of 15–20 % at sotalol concentrations of 1.8–14 μ M (Braam et al. 2010). In a subsequent study, they also observed 20 % cFPD prolongation in hESC-CMs at sotalol concentration of 10 μ M (Braam et al. 2013). Our study resulted in similar data. At concentrations of 6–10 μ M, we observed a cFPD prolongation of 12–18 % in the hESC-CMs, increasing to 21 % at 19 μ M. We observed that sotalol increased cFPD in all the CMs at the therapeutical concentration range (1.6–9.7 μ M). In line with our data, Mehta et al. (2013) reported similar observations from sotalol-induced cFPD prolongation in hiPSCderived CMs (Mehta et al. 2013). Furthermore, according to our data sotalol significantly increased cFPD at earlier concentrations in LQT-CMs than in WTa or WTb. This may suggest that LQT-CMs may be more vulnerable to the cFPD-prolonging effect of sotalol than hiPSC-derived control cells.

Quinidine is a class Ia antiarrhythmic drug that primarily blocks sodium ion channels at high concentrations and hERG channel at low concentrations and is associated with TdP (Bauman et al. 1984). Quinidine prolonged the cFPDs in all of our cell lines ranging from 16 to 42 %. Braam et al. (2010) reported a prolongation of 20-50 % in hESC-CMs with quinidine concentrations of up to 3.2 μ M and increases in the cFPD at higher concentrations, and Caspi et al. (2009) reported a cFPD







prolongation of slightly <30 % in hESC-CMs with 8 μ M quinidine (Braam et al. 2010; Caspi et al. 2009). Mehta et al. (2013) reported a 20 and 55 % cFPD prolongation in 150–170 days old hiPSC-CMs at 1 and 10 μ M concentrations, respectively. Here, we noticed an 18 % increase in the cFPD of the hESC-CMs at a concentration of 9 μ M. Quinidine induced beating arrests which emerged at the

average or upper therapeutic serum concentration. Quinidine is known to inhibit sodium ion channels at high concentrations (Snyders and Hondeghem 1990), which may be the reason for higher frequency of beating arrest than AD-like waveform abnormalities. The LQT2-CMs, showed significantly prolonged cFPD already at the half of the lower limit of therapeutic concentration (1.5 μ M),

Sotalol (µM)	hESC (n = 11)	WTa (n = 6)	WTb (n = 12)	LQT1A (n = 13)* ^{,s}	LQT1B (n = 13) ^{*,a}	LQT2 (n = 15) ^a
0.8						
1.6						
5.7						AD(2)
9.7	AD(4)		AD(1)			
19.4		AD(1)		AD(2)		
Quinidine (µM)	hESC (n = 10)	WTa (n = 6)	WTb $(n = 11)$	LQT1A (n = 12)*, ^s	LQT1B (n = 16)* ^{,a}	LQT2 $(n = 12)^{a}$
1.5						
3						AD(1)
9	AD(3)		BA(7)	AD(1)	BA(5)	BA(5)
15		BA(2)		BA(2)		
30	BA(1)					
E-4031 (nM)	hESC (n = 10)	WTa (n = 6)	WTb (n = 8)	LQT1A (n = 16)*, ^s	LQT1B (n = 16)* ^{,a}	LQT2 $(n = 13)^{a}$
10						
100						BA(4)
300	AD(7)	AD(2)		AD(8)		AD(3)
500	BA(1)	BA(2)	BA(1)		AD(4)	
700					BA(1)	

Table 3 Arrhythmic effects of sotalol, quinidine, and E-4031 observed during the field potential (FP) recordings

The reported values in the table are the lowest concentrations that induced the effect. The numbers in parentheses indicate how many cardiomyocyte clusters in total were affected

AD afterdepolarization (AD)-waveforms, BA beating arrest (termination of FP activity)

* Siblings ^s symptomatic ^a asymptomatic

unlike control- or LQT1-CMs, and AD-like waveform abnormalities at the lower limit of therapeutic concentration level (3 μ M). These findings together may imply that LQT2 harboring hERG R176 W may be at more increased risk for quinidine-induced cFPD prolongation and proarrhythmic effects than control- or LQT1-CMs harboring G589D mutation. However, we emphasize that our findings, based on cell lines derived from three LQTS patients only, must be interpreted with caution.

E-4031 is a hERG channel blocker (Caspi et al. 2009) that reduced the I_{Kr} current during cardiac repolarization and caused prolongations in all our cell lines. Braam et al. (2010) reported FPD prolongation in hESC-CMs treated with 30 nM E-4031 and AD-like waveform abnormalities at concentrations of $1 \mu M$ and over (Braam et al. 2010). In hiPSC-CM clusters similar cFPD prolongations have been reported as well (Mehta et al. 2011, 2014). We observed that the cFPD increased significantly more in CMs derived from symptomatic LQT1 patient than from asymptomatic. Presently, it is not known whether this indicates that symptomatic LQT1 may be at more increased risk for hERG blocker-induced cFPD increase. Egashira et al. (2012) reported cFPD prolongation of 20 % in LQT1-CMs derived from a symptomatic patient, when treated with 100 nM E-4031. These LQT1-CMs had a heterozygous deletion mutation in KCNQ1, 1893delC (Egashira et al. 2012). Here, we found cFPD prolongation of 13 and 8 % in LQT1A- and LQT1B-CMs carrying a G589D missense mutation. Egashira et al. (2012) also reported AD-like waveform abnormalities in LQT1-CMs treated with 300 nM E-4031, which is in line with our results from symptomatic LQT1A-CMs.

Variations of the in vitro responses to QTc interval affecting drugs may result from several reasons. Differences in the common polymorphisms regulating cardiac repolarization may result in the modulation of drug responses (Roden 2004). Furthermore, differences in drug responses between the LQT1 and -2 subtypes may result from differences in their respective repolarization reserves (Roden 1998). Other possible explanations may come from the observations that hPSC-CMs can have very low or very high spontaneous beating rates, and that the hPSC-CMs with $BR \ge 50$ bpm express more atrial-related genes and therefore might be more atrial-like (Asp et al. 2010). Different cardiomyocyte subtypes (e.g., atrial vs. ventricular) may have different ion channel compositions (Gaborit et al. 2007). Therefore, for certain applications, if the beating rates are representative of differences between cardiac subtypes, then the drug responses might vary accordingly. We did not test whether the END-2 differentiation protocol used in this study generated more nodal-, atrial- or ventricular-like CMs across the cell lines. The MEA analysis used

in the present study includes only beating clusters, suggesting the presence of nodal-, atrial- and ventricular-like type of CMs. However, previous study has shown that END-2 differentiation protocol generates mostly ventricular CMs shown by electrophysiological measurements (Mummery et al. 2003). It is also unknown whether the heterozygous mutated ion channels are assembled similarly in all differentiating cardiomyocytes. Variation in the ion channel composition from cluster to cluster and even from cell to cell might lead to differential drug responses. There are some evidence that there may be differences in same certain drug responses between clusters of CMs and monolayer CMs. In disease-free hiPSC-CM monolayers, Harris et al. (2013) reported statistically significant E-4031 induced cFPD prolongation already at 3 nM whereas for disease-free clusters of hESC- or hiPSC-CMs the significant cFPD prolongations were reported to be found in 30-300 nM (Braam et al. 2010; Caspi et al. 2009; Mehta et al. 2011, 2014). In diseasefree hiPSC-CM monolayers E-4031 evoked arrhythmias were seen as early as 10-30 nM (Harris et al. 2013; Nakamura et al. 2014). We have also observed similar phenomenon in monolayer-based disease-free CMs with E-4031 (data not published). The aforementioned differences may have significant implications in drug screenings and would require further investigation.

Potential limitation of the study

The clinical phenotypes of the patients did not correlate with repolarization times in this study. The occurrence of clinical symptoms in LQT-patients harboring G589D have been linked to QTc value. In the group of QTc > 500 ms over 50 % of the patients were symptomatic whereas only 16 % were symptomatic with QTc < 440 ms (Piippo et al. 2001). In addition, patients with C-terminal mutation such as G589D have milder phenotypes compared to transmembrane domain mutations of the same channel (Donger et al. 1997; Larsen et al. 1999; Piippo et al. 2001; Swan et al. 1999). In this study, the LQT1-G589D-patients' QTc values were 456 and 428 ms for symptomatic and asymptomatic, respectively. In LQT2, R176W mutation has been reported elsewhere as a relatively mild form with modest QTc prolongation (Fodstad et al. 2006; Marjamaa et al. 2009). Here, the LQT2 patient had been asymptomatic except for occasional palpitations. In the present study, at the baseline significant differences in repolarization times were observed, but the drug effects appeared to be of similar magnitude in LQT-CMs harboring G589D or R176W mutations compared to healthy controls at this sample size. Given the large variance in cFPD it may be possible that by increasing the sample size statistical significances could be found between control and LQT-CMs. However, it should be emphasized that increases in repolarization times which are already at baseline longer are potentially more severe that in those of normal length.

Conclusion

In conclusion, the present study reports in vitro electrophysiological effects of various non-cardiac and cardioactive drugs as assessed in hiPSC-derived LQTcardiomyocytes. We have shown that long QT syndrome is recapitulated in hiPSC-derived LQT-cardiomyocytes. All drugs tested showed a systematic tendency to prolong the field-potential duration (cFPD, an indicator of repolarization phase). We did not observe any systematic differences in the sensitivities of cFPD prolongation among different drugs between control- and LQT-cardiomyocytes. Further studies are needed to explore the possibility that this system could aid in the patient-specific risk assessment when considering the use of potentially QTinterval prolonging drugs.

Additional file

Additional file 1: Figure S1. Immunocytochemical marker expression of human induced pluripotent stem cells (hiPSCs) and their cardiomyocyte derivatives. **Table S1.** Pluripotency marker expression of the hPSCs. (n.d.: not determined).

Authors' contributions

JK performed MEA experiments, analyzed MEA data and wrote the manuscript, VJK designed and performed MEA experiments, analyzed MEA data and wrote the manuscript, AK generated and characterized iPS cell lines, performed MEA experiments and wrote parts of the manuscript, MO generated and characterized control iPS cell line and wrote parts of the manuscript, HS participated to the conception of the study and revised the manuscript critically for important intellectual content, KK participated to the conception of the study and revised the manuscript critically for important intellectual content, KA-S conceived the study, designed experiments and wrote the manuscript. All authors read and approved the final manuscript.

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Acknowledgements

We thank Henna Venäläinen and Merja Lehtinen for excellent technical help. We also thank Mika Helminen for performing statistical analyses and Paruthi Pradhapan for providing support to CardioMDA software. The END-2 cells were a kind gift from Professor Christine Mummery. This study was funded by grants from the Academy of Finland, the Finnish Cultural Foundation, Biocenter Finland, the Finnish Foundation for Cardiovascular Research, Pirkanmaa Hospital district funding (EVO), the Foundation of Aarne Koskelo and the Finnish Funding Agency for Technology and Innovation (TEKES).

Competing interests

The authors declare that they have no competing interests.

Received: 25 November 2015 Accepted: 17 February 2016 Published online: 29 February 2016

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