Modeling Short QT Syndrome Using Human-Induced Pluripotent Stem Cell–Derived Cardiomyocytes

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Background—Short QT syndrome (SQTS), a disorder associated with characteristic ECG QT-segment abbreviation, predisposes affected patients to sudden cardiac death. Despite some progress in assessing the organ-level pathophysiology and genetic changes of the disorder, the understanding of the human cellular phenotype and discovering of an optimal therapy has lagged because of a lack of appropriate human cellular models of the disorder. The objective of this study was to establish a cellular model of SQTS using human-induced pluripotent stem cell–derived cardiomyocytes (hiPSC-CMs).

Methods and Results—This study recruited 1 patient with short QT syndrome type 1 carrying a mutation (N588K) in KCNH2 as well as 2 healthy control subjects. We generated hiPSCs from their skin fibroblasts, and differentiated hiPSCs into cardiomyocytes (hiPSC-CMs) for physiological and pharmacological studies. The hiPSC-CMs from the patient showed increased rapidly activating delayed rectifier potassium channel current (IKr) density and shortened action potential duration compared with healthy control hiPSC-CMs. Furthermore, they demonstrated abnormal calcium transients and rhythmic activities. Carbachol increased the arrhythmic events in SQTS but not in control cells. Gene and protein expression profiling showed increased KCNH2 expression in SQTS cells. Quinidine but not sotalol or metoprolol prolonged the action potential duration and abolished arrhythmic activity induced by carbachol.

Conclusions—Patient-specific hiPSC-CMs are able to recapitulate single-cell phenotype features of SQTS and provide novel opportunities to further elucidate the cellular disease mechanism and test drug effects. (J Am Heart Assoc. 2018;7:e007394. DOI: 10.1161/JAHA.117.007394.)

Key Words: arrhythmia (heart rhythm disorders) • arrhythmia (mechanisms) • ion channel • short QT syndrome

It has been reported that an alteration of QTc interval over 24 hours, either shortening or prolongation, is associated with sudden cardiac death (SCD).1 Short QT syndrome (SQTS) is a rare, inheritable cardiac electrical disease characterized by a shortened corrected QT interval (QTc) and associated with a risk of SCD. The prevalence of SQTS is 0.02% to 0.1% with a male predominance.2 The majority of cardiac events (>80%) in SQTS patients occur under resting conditions or during sleep. Even survivors of SCD are at high risk for recurrent ventricular tachyarrhythmia events. So SCD remains...
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Clinical Perspective

What Is New?

- This is the first cellular model of short QT-syndrome type 1 using human induced pluripotent stem cell–derived cardiomyocytes.
- The site mutation (p.N588K) in KCNH2 increases the expression of HERG channels and thus enhances outward potassium current, shortens action potential duration, and promotes proarhythmic activity in short QT-syndrome–cardiomyocytes.

What Are the Clinical Implications?

- Patient-specific pluripotent stem cell–derived cardiomyocytes may provide opportunities to gain new insights into the cellular mechanisms of short QT-syndrome and discover new effective drugs for treating the disease.

a major public health burden despite revolutionary progress in the past 3 decades in the treatment of ventricular tachyarrhythmia with the use of implantable cardioverter defibrillator (ICD) therapy.

The QT interval of the human ECG presents the duration of the cellular action potential. It depends on the heart rate and therefore a correction should be made before its interpretation. The latest European Society of Cardiology guideline produced in 2015 suggests upper and lower limits of 480 and 360 ms, respectively, for both males and females.3 The presence of QTc ≤2015 suggests upper and lower limits of 480 and 360 ms, respectively, for both males and females. The QT interval of the human ECG presents the duration of the cellular action potential. It depends on the heart rate and therefore a correction should be made before its interpretation. The latest European Society of Cardiology guideline produced in 2015 suggests upper and lower limits of 480 and 360 ms, respectively, for both males and females.3 The presence of QTc ≤2015 suggests upper and lower limits of 480 and 360 ms, respectively, for both males and females.3 The presence of QTc ≤2015 suggests upper and lower limits of 480 and 360 ms, respectively, for both males and females.3 The presence of QTc ≤2015 suggests upper and lower limits of 480 and 360 ms, respectively, for both males and females.3 Therefore, we aimed in the present study to establish a cellular model of short QT-syndrome and discover new effective drugs for treating the disease.

SQTS can have a congenital cause or acquired causes. The first family with the entity of SQTS was described in 2000.8 After that, more patients with SQTS have been reported.4,9,10 Until now, 6 subtypes of SQTS are known. SQTS1, SQTS2, and SQTS3 are associated with gain of function of potassium channels, whereas SQTS4-6 can be caused by loss of function of calcium channels. The first genetic alteration in SQTS was identified in the KCNH2 gene encoding a pore-forming subunit of voltage-activated potassium channel belonging to the ether-a-go-go family, which mediates the rapidly delayed rectifying potassium current (IKr) in the heart. Currently, 3 genes encoding potassium channels, KCNQ2 (for SQTS1), KCNQ1 (for SQTS2), KCNJ2 (for SQTS3), and 3 more genes encoding calcium channels, CACNA1C (for SQTS4), CACNB2 (for SQTS5), and CACNA2D1 (for SQTS6) are identified and linked to SQTS. The genetic analysis, however, does not identify any genetic cause in up to 40% of SQTS cases.11 Additionally, overlap syndrome, SQTS with Brugada syndrome, has been reported.12,13 Causes of acquired SQTS are electrolyte imbalance including hyperkalemia or hypercalcemia, myocardial ischemia, acidosis, or carnitine deficiency. Hyperthermia can also cause a shortened QT interval, as can drugs such as digitalis, acetylcholine, catecholamines, or ATP-sensitive K⁺ channel activators. Short QT intervals have also been associated with epilepsy, particularly during the ictal and postictal states.14–16

In spite of rapid advances in understanding the mechanisms and genetic bases of SQTS, much less is known about the appropriate treatments for SQTS patients. Because of the high risk of SCD, it has been recommended to implant ICD in patients with a strong family history of SCD and evidence of short QTc interval. However, ICD is not always feasible or adequate for every patient. Therefore, effective drug therapies are needed for some patients. Until now, there has been very limited clinical research to test drug effects in SQTS patients. A few antiarrhythmic drugs such as disopyramide, nifekalant, quinidine, flecainide, sotalol, and ibutilide have been tested in small patient series with SQTS1, but only quinidine has been shown to be effective in the treatment.10,17–19

For studies on cardiac functions, especially the cardiac ion channel functions, human induced pluripotent stem cell–derived cardiomyocytes (hiPSC-CMs) have important advantages over heterologous expression systems such as Xenopus oocytes, human embryonic kidney cells, and Chinese Hamster Ovary cells lacking important constituents of cardiac ion channel macromolecular complexes that might be necessary for normal electrophysiological characteristics. Transgenic animals possess cardiac electrophysiological properties crucially different from that in humans. Thus, taking into account the hurdle for obtaining human ventricular cardiomyocytes, hiPSC-CMs could be a good alternative for SQTS studies, either mechanistic or therapeutic. Indeed, hiPSC-CMs have been successfully used to recapitulate the phenotype of some genetic heart diseases such as long QT syndromes, Brugada syndrome, arrhythmogenic right ventricular cardiomyopathy, catecholaminergic polymorphic ventricular tachycardia, and hypertrophic and dilated cardiomyopathy.20–24

Therefore, we aimed in the present study to establish a cellular model of SQTS, which is still lacking, by using hiPSC-CMs from a short QT patient with a mutation (N588K) in the KCNH2 gene.

Materials and Methods

The data, analytic methods, and study materials will be made available to other researchers for purposes of reproducing the
results or replicating the procedure; they will be provided on request to the corresponding author.

Ethics Statement

The skin biopsies from 2 healthy donors and 1 SQT patient were obtained with written informed consent. The study was approved by the Ethics Committee of the Medical Faculty Mannheim, University of Heidelberg (approval number: 2009-350N-MA) and by the Ethics Committee of University Medical Center Göttingen (approval number: 10/9/15). The study was carried out in accordance with the approved guidelines and conducted in accordance with the Helsinki Declaration of 1975, as revised in 1983.

Clinical Data

Skin biopsy was performed in a 29-year-old male patient with familial SQTS1 caused by identified missense mutation (C to G substitution at nucleotide 1764) in the KCNH2 gene. The SQTS was diagnosed many years ago. Two patients from a family were referred for syncope, palpitations, and a strong family history of SCD, which was present in 3 generations. The family pedigree is demonstrated in Figure 1A. Three of the family members including our patient presented a short QT interval (270–300 ms corrected with Bazett’s formula (QTc) at ECG (Figure 1B). Extensive clinical and instrumental evaluation, including physical examination, serial ECGs, 24-hour Holter monitoring, echocardiogram, exercise testing, and cardiac magnetic resonance imaging confirmed familial SQTS. The electrophysiology study of this patient showed a ventricular effective refractory period of 140 ms at any pacing site and at different pacing cycle lengths. Programmed stimulation did not induce malignant arrhythmia. Atrial effective refractory period was 160 ms at a basal cycle length of 600 ms. In consideration of the familial history and presence of SQT, the patient prophylactically received an ICD. Nineteen months after implantation the patient had an appropriate ICD shock because of a documented ventricular fibrillation.

Generation of Human iPS Cells

Human iPSC cells (hiPSCs) were generated from primary human fibroblasts derived from skin biopsies. HiPSC line D1 (here abbreviated as D1) was generated using lentiviral particles carrying the transactivator rtTA and an inducible polycistronic cassette containing the reprogramming factors OCT4, SOX2, KLF4, and c-MYC as described previously.27 HiPSC lines isSTQSa1.7 (GOEi091-A.7, here abbreviated as SQTS), isSQTSa1.8 (GOEi091-A.8), and isSQTSa1.15 (GOEi091-A.15) were generated in feeder-free culture conditions using the integration-free CytoTune-iPS 2.0 Sendai Reprogramming Kit (Thermo Fisher Scientific, #A16517) with the reprogramming factors OCT4, KLF4, SOX2, and c-MYC according to manufacturer’s instructions with modifications. The generated hiPSCs were characterized for their pluripotency and their in vitro differentiation potential (Figures S1 and S2). See Data S1 for further details.

Polymerase-Chain-Reaction Assays

To quantify the steady-state mRNA expression of the hiPSC-CMs, RNA was reverse transcribed and quantitative polymerase chain reaction was performed as described.29 Gene symbols, RefSeq No., and Cat. No. of the primers used for quantitative polymerase chain reaction analyses in hiPSC-CMs characterization are listed in Table S1. For evaluation of the characteristics of the used hiPSC lines, reverse transcription-polymerase chain reaction was performed as follows: Total RNA was isolated using the SV Total RNA Isolation System (Promega, #Z3105) according to manufacturer’s instructions. One hundred nanograms RNA was used for the first-strand cDNA synthesis by using MULV Reverse Transcriptase (Thermo Fisher Scientific, #N8080018) and Oligo d(T)16 (Thermo Fisher Scientific, #N8080128). One tenth of cDNA was used as polymerase chain reaction template and amplified using the GoTaq G2 DNA polymerase (Promega, #M7845) according to manufacturer’s instructions. Primer sequences, annealing temperatures, and cycles used for reverse transcription polymerase chain reaction analyses of the hiPSC lines are listed in Table S2.

Immunofluorescence Staining

Immunofluorescence staining was performed using appropriate primary antibodies and AlexaFluor conjugated secondary
antibodies (ThermoFisher) according to the manufacturer’s instructions. The primary antibodies used in hiPSC-CMs were α-actinin (Sigma Aldrich), TNNT (Sigma Aldrich), and KCNH2 (Sigma Aldrich). All the other antibodies used for characterization of hiPSC lines are listed in Table S3.

Patch-Clamp

Standard patch-clamp recording techniques were used to measure the rapidly delayed rectifier potassium (I_{Ko}), slowly delayed rectifier potassium (I_{Ks}), transient outward potassium (I_{to}), inward rectifier (I_{ki}), late sodium (I_{Na}), and L-type calcium (I_{CaL}) channel currents as well as action potential (AP) in the whole-cell configuration at room temperature.

Measurement of Intracellular Calcium Concentration

To measure the intracellular Ca^{2+} concentration ([Ca^{2+}]), cells were loaded with the fluorescent Ca^{2+}-indicator Fluo-3 AM. The fluorescence of the cells was measured by using Cairn Optoscan calcium imaging system (Cairn Research, UK). Fluorescence is excited by 488 nm and emitted at 520 nm. Changes in [Ca^{2+}] were described by

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Figure 1. A, The pedigree of the family with SQTS. The patient recruited for this study is marked by the arrow. B, The ECG from the SQTS patient shows a classic SQTS-ECG pattern. SCD indicates sudden cardiac death; SQT, short QT syndrome (SQTS).
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[Ca^{2+}] = k_d \left( \frac{F}{F_{\text{max}}} - F \right)
\]

where \(k_d\) = dissociation constant of Fluo-3 (400-nmol/L), \(F\) = Fluo-3 fluorescence, and \(F_{\text{max}} = \text{Ca}^{2+}\)-saturated fluorescence obtained at the end of each experiment.\(^{30}\)

**Statistical Analysis**

If not otherwise indicated, data are shown as mean±SEM and were analyzed using InStat© (GraphPad, San Diego, CA) and SigmaPlot 11.0 (Systat GmbH, Germany). By analyzing the data with the Kolmogorov Smirnov test, it was decided whether parametric or nonparametric tests were used for analysis. Student t test and the Mann–Whitney U test were used to compare continuous variables with normal and non-normal distributions, respectively. The \(\chi^2\) test for independence was used to compare categorical variables. For parametric data, 1-way ANOVA with Bonferroni post-test for multiple comparisons was performed. For nonparametric data, the Kruskal–Wallis test with Dunn’s multiple comparisons post-test was used. Paired t test was used for comparisons of data before and after application of a drug. \(P<0.05\) (2-tailed) was considered significant.

**Results**

**Characterization of Patient-Specific hiPSCs and hiPSC-CMs**

To confirm the successful generation of hiPSCs reprogrammed from skin fibroblasts of a patient with SQTS and a healthy donor, 3 iPSC lines of each subject were first characterized for pluripotency. Generated iPSCs displayed characteristic human embryonic stem cell morphology and showed expression of pluripotency markers at mRNA and protein level (Figure 2A through 2D, Figure S1A through S1C and S2A through S2C). Spontaneous differentiation of generated iPSC lines was analyzed by embryoid body formation and germ-layer specific marker expression. Immunocytochemical staining of spontaneously differentiated hiPSC lines shows expression of endodermal marker AFP (α-fetoprotein), mesodermal-specific α-SMA (α-smooth muscle actin), and ectodermal βIII-tubulin. Nuclei are co-stained with DAPI. Scale bars: 100 μm.

**Figure 2.** Pluripotent characteristics of human induced pluripotent stem cell (hiPSC) lines. A, The hiPSC lines of short QT syndrome (SQTS) (SQTSa1.7) and donor 2 (D2) (ipWT1.3) generated from skin fibroblasts (upper panel) display a typical morphology for human pluripotent stem cells (lower panel). B, In comparison to donor’s fibroblasts, generated iPSC lines show expression of endogenous pluripotency markers SOX2 (sex determining region Y- box 2), OCT4 (octamer-binding transcription factor 4), NANOG (pron. nanOg, homeobox protein), LIN28 (lin-28 homolog A), FOXD3 (Forkhead Box D3), and GDF3 (growth differentiation-factor-3) at mRNA level proven by reverse transcription-polymerase chain reaction. Human embryonic stem cells (hESCs) were used as positive control, and mouse embryonic fibroblasts (MEFs) were used as negative control. C, Generated hiPSC lines express pluripotency markers OCT4, SOX2, NANOG, LIN28, SSEA4 (stage-specific embryonic antigen 4), and TRA-1-60 as shown by immunofluorescence staining. Nuclei are co-stained with DAPI (4’, 6-diamidino-2-phenylindole). D, Flow cytometry analysis of pluripotency markers OCT4 and TRA-1-60 reveals a homogeneous population of pluripotent cells in generated iPSC lines. E, Spontaneous differentiation potential of generated hiPSC lines was analyzed by embryoid body formation and germ-layer specific marker expression. Immunocytochemical staining of spontaneously differentiated hiPSC lines shows expression of endodermal marker AFP (α-fetoprotein), mesodermal-specific α-SMA (α-smooth muscle actin), and ectodermal βIII-tubulin. Nuclei are co-stained with DAPI. Scale bars: 100 μm.
conditions for subsequent functional analyses. Beating cardiomyocytes were observed 8 to 12 days after starting the differentiation. No differences in differentiation efficiency were observed between control and SQTS iPSCs. Typical cardiac markers such as α-actinin and human cardiac troponin T were presented at day 40 of differentiation (Figure 3A and 3B). The beating cardiomyocytes 40 to 50 days after differentiation were used for patch-clamp and calcium transient imaging.

**Action Potential Duration Was Shortened in hiPSC-CMs From the Patient**

AP characterizations are summarized in Table S4. Action potential amplitude, resting potential, and maximum depolarization velocity ($V_{\text{max}}$) were similar in the measured hiPSC-CMs from the patient and healthy donors (Figure 4B through 4D). Action potential durations (APD 50 and 90), however, were significantly shorter in SQTS cells compared with that in the cells from the 2 healthy donors (Figure 4E and 4F). Of note, cells from the 2 donors displayed very similar AP parameters.

**$I_{\text{Kr}}$ was Enhanced in hiPSC-CMs From the Patients**

To investigate the underlying currents for APD shortening, we assessed different ion channel currents that can influence APD. First, we used patch-clamp technique and specific ion channel blockers to measure and compare the $K^+$ currents in SQTS cells and donor cells. The transient outward current ($I_{\text{to}}$), the rapidly activating delayed rectifier potassium current ($I_{\text{kr}}$) and slowly activating delayed rectifier potassium current ($I_{\text{ks}}$)
as well as inward rectifier potassium channel current (I_{K1}) were assessed by 4-AP-, E-4031-, chromanol 293B-, and Ba^{2+}-sensitive currents, respectively. As expected, I_{Kr} was enhanced in SQTS cells, but I_{to} and I_{ks} as well as I_{K1} were similar among all the cells (Figures 5 and 6, Figures S3 and S4). Notably, the steady-state currents of I_{Kr} were enhanced but the tail currents were reduced in SQTS cells (Figure 5C), indicating an impairment of inactivation of KCNH2 channels. In addition, we also checked L-type calcium channel current (I_{CaL}) and late sodium channel current (I_{NaL}), which can also influence APD. Both currents were not significantly changed in SQTS cells when compared with donor cells (Figure 6D and 6E, Figure S5).

Considering that steady-state inward currents were always observed in all the cell lines and all the depolarizing pulses induced instantaneous peak currents when the holding potential was set at −40 mV, which are different from that observed in a previous study, some I_{Kr} recordings were performed at a holding potential of −80 mV. When the holding potential was changed to −80 mV, the basal inward currents disappeared and the instantaneous currents were replaced by currents showing time-dependent activation (Figure S6).

Next, by using quantitative polymerase chain reaction analysis, we evaluated the gene expression of the assessed ion channels. The mRNA levels of KCND3/KCHIP2 (for I_{to}), CACNA1C (for α-subunit of I_{CaL}), and KCNH2 (for I_{Kr}) were significantly increased in SQTS-hiPSC-CMs, whereas the gene expression of SCN5A (for I_{NaL}), KCNJ2 (for I_{K1}), and of KCNQ1 (for I_{ks}) were similar compared with donor hiPSC-CMs.
The quantitative polymerase chain reaction data except for CACNA1C and KCND3/KCHIP2 were in agreement with patch-clamp data. More strikingly, besides enhanced current and mRNA level, the increased protein level of KCNH2 channels in the cell membrane was also detected by the immunostaining of KCNH2 (Figure 3E through 3H).

Calcium Handling Was Changed and Arrhythmia Events Were Increased in hiPSC-CMs From the Patients

To check whether the calcium handling is normal in SQTS-hiPSC-CMs, we examined the intracellular calcium concentration. An increasing of systolic and diastolic concentration has been observed (Figure 8A through 8E). These results reveal altered Ca^{2+} homeostasis in human SQTS cardiomyocytes with mutation in the HERG-gene.

Interestingly, during measurements of spontaneous calcium transient, all SQTS cardiomyocytes presented irregular, delayed afterdepolarization–like and early afterdepolarization–like arrhythmic events, whereas many fewer donor cells showed those phenomena. The percentage of cells showing arrhythmic events in SQTS, D1, and D2 cells are 100%, 50%, and 37.5%, respectively ($P<0.05$, Figure 8F). Given that arrhythmias in SQTS patients happen frequently at rest or in sleep, we checked whether cholinergic stimulation can be a trigger for arrhythmias. Indeed, 10 μmol/L carbachol (CCh), a muscarinic receptor agonist, provoked arrhythmic events in cardiomyocytes from the SQTS patient (percentage of cells showing CCh-induced arrhythmic events: SQT, 100%; D1, 0%; D2, 14%; $P<0.05$; Figure 9C).

Quinidine Attenuated Arrhythmic Events in hiPSC-CMs From the Patients

To test the responses of the SQTS-hiPSC-CMs to antiarrhythmic drugs, we chose 3 drugs, quinidine, sotalol, and metoprolol, which have been assessed in patients or cells transfected with the same mutated $I_{Kr}$ channels in previous studies. All 3 drugs did not change resting potential and action potential amplitude (Figure S7). Ten micromoles per liter quinidine reduced $V_{\text{max}}$ (Figure S7A), prolonged APD (Figure S8A through S8C), and also abolished CCh-induced arrhythmic events in SQTS cardiomyocytes (Figure 9D).
through 9F), while sotalol and metoprolol were not effective in prolonging APD (Figure S8D through S8F, Table S4), consistent with previous in vitro and in vivo studies. Taken together, our data indicate that our hiPSC-CMs from the SQTS patient mimicked the phenotype of SQTS and recapitulated the drug effects in SQTS patients.

Discussion

We have for the first time generated patient-specific induced pluripotent stem cell–derived cardiomyocytes from a SQTS patient with “aborted sudden cardiac death” carrying a mutation in the KCNH2 gene and compared their cellular physiological and pharmacological properties with that of healthy cells. We have found the following in SQTS-hiPSC-CMs: (1) a shortening of APD and enhancement of Ikr; (2) elevated expression level of KCNH2 channels; (3) a high vulnerability of SQTS cardiomyocytes with arrhythmia events; (4) carbachol-provoked arrhythmia events in SQTS cardiomyocytes; and (5) the antiarrhythmic drug quinidine but not metoprolol or sotalol prolonged APD and attenuated arrhythmia events.
SQTS is a newly recognized rare cardiac disorder. This disease has not been well investigated yet. Many data, either clinical or experimental, which are important for understanding or treating the disease, are not available. One obstacle for studies on SQTS is the low prevalence. It is difficult to have enough patients or tissues from patients for mechanistic or drug-testing studies. Therefore, an optimal experimental model for SQTS will be useful for studying this rare disease. Recently, an animal model of drug-induced SQTS has been established to test candidate antiarrhythmic agents that may be effective for treatment of SQTS patients.32–34 Because of the differences between animals and humans, animal models are not ideal for drug testing. In the current study, we used patient-specific hiPSC-CMs to model SQTS and test drug effects.

The patient chosen for this study is from a family with familial SQTS1 carrying a missense mutation (C to G substitution at nucleotide 1764) in KCNH2 (HERG/IKr) gene, which results in an amino acid change in the S5-P loop region (p.N588K) in HERG/IKr channels. The clinical characteristics of the family including syndromes, family history, gene investigation, ECG properties, and treatment have been described in detail in previous publications.5,19,35 Importantly, all the family members showed SQTS or sudden cardiac arrest but none of the unaffected members in the family carry the mutation.35 So it is already clear that this mutation is associated with SQTS in this family. The influence of this mutation on the channel current (IKr) has been experimentally investigated.35–38 Effects of some antiarrhythmic drugs have also been tested in this family.38 All the published data can be useful for determining the success of the planned modeling of the SQTS phenotype by using hiPSC-CMs. The first aim of this study was to check whether the hiPSC-CMs from a SQTS patient of this family can model the disease. Indeed, the hiPSC-CMs from the patient of the family displayed enhanced IKr, abbreviation of APD, and increased arrhythmia events.

**Figure 6.** Changes in ion channel currents in SQTS (short QT syndrome)-cardiomyocytes. Ion channel currents were recorded in the control (D1, D2) and SQTS-cells (SQTS). A, Mean values of I_{Ks} (slowly activating delayed rectifier potassium channel). B, Mean values of peak I_{to} (transient outward potassium channel current). C, Mean values of I_{K1} (inward rectifier potassium channel current). D, Mean values of peak I_{CaL} (L-type calcium channel current). E, Mean values of the area under curve of TTX (tetrodotoxin) sensitive late I_{Na} (sodium channel current). Values given are mean±SEM; n, number of cells; ns (not significant), P>0.05.

DOI: 10.1161/JAHA.117.007394
indicating successful recapitulation of the main phenotype of SQTS in the patient.

Regarding the mechanism underlying the shortening of QT-interval in the family, the HERG channel was mutated at the same site as in the patients (p.N588K-HERG) and was expressed in modified human embryonic kidney cells (TSA201) to check the effects of the mutation on channel current. Interestingly, it was revealed that the p.N588K-mutation enhanced the channel currents (Ikr), suggesting that gain of function mutation of HERG channel, which is an important repolarizing current and determinant for APD, may be the reason for the abbreviation of QT-interval. Also, it was proved that the p.N558K-mutation led to impairment of channel inactivation and thus enhanced the steady-state currents, representing the process of the channel activation from resting state to open state (Figure S6). Of note, the enhanced outward steady-state currents and reduced tail currents in SQTS cells could be observed when the holding potential was set at either −40 mV or −80 mV.

Tachyarrhythmias in SQTS patients occur frequently at rest or in sleep, suggesting involvement of increased vagal tone. A previous study showed that acetylcholine applied to coronary-perfused canine left-ventricular wedge preparations, mimicking increased vagal tone, precipitated drug- (an IcaL activator plus a ICaL blocker) induced polymorphic ventricular tachycardia/ventricular fibrillation. Until now, this effect has not been tested in a SQTS model. We observed in SQTS-hiPSC-CMs that a muscarinic receptor agonist, CCh, increased arrhythmia events compared with cells from 2 healthy donors. This finding supports the role played by vagal tone in occurrence of tachyarrhythmias in SQTS patients. Given that CCh alone did not show any effect on INa, ICaL, IKs, IKr, and IKs, the mechanisms underlying the CCh-induced arrhythmia events need to be clarified in future studies.

For the treatment of SQTS, there is still no optimal strategy. Because of SCD risk, ICD has been recommended and is also effective in some patients. Because of the limitations of ICD and to avoid inappropriate ICD shock delivery that is also associated with increased mortality, drug therapy is also needed. Therefore, different antiarrhythmic drugs were tested in different SQTS families including our patient. Surprisingly, the well-known IKr blocker, sotalol, did not effectively prolong QT-interval, whereas quinidine, a multiple-channel blocker, corrected QT-abbreviation and prevented arrhythmias. The reason for ineffectiveness of sotalol has been proved in a heterologous expression system (TSA201 cells) to be the largely reduced sensitivity of the mutated channel to sotalol. The second aim of the present study was to test whether the hiPSC-CMs from the SQTS patient exhibit the same or similar responses to the examined drugs. Actually, we observed in SQTS-hiPSC-CMs that sotalol failed but quinidine succeeded in prolonging APD.
and reversed CCh-induced arrhythmia events, again in good agreement with previous data of either clinical or experimental studies. This finding highlighted the ability of iPSC-CMs to recapitulate not only the phenotype but also the drug responses of the disorder. Furthermore, another study tested the effects of 2 widely used β-blockers, carvedilol and metoprolol, on p.N588K-HERG and p.V307L-KCNQ1 channels expressed in mammalian cells. It was found that metoprolol is more potent than carvedilol in inhibiting both mutant K+ channel currents. This suggests that metoprolol may be a new candidate drug for treating SQTS. However, in the current study, metoprolol failed to prolong APD in SQTS-hiPSC-CMs. This suggests effects of metoprolol on other ion channels that counteracted the Ikr and Ifs inhibition. Thus, our data argue against the application of metoprolol for SQTS treatment.

**Conclusion**

Our data demonstrate that the main phenotypic features of SQTS were successfully recapitulated in the hiPSC-CMs from a SQTS1 patient, including increased Ikr, shortened APD, and increased arrhythmia events as well as changed responses to antiarrhythmic drugs. In addition, a new mechanism for Ikr enhancement in SQTS1 with p.N588K mutation was identified. This study supports an application of hiPSC-CMs in future studies on SQTS including examination of disease mechanisms, identification of new therapeutic targets, and test of drug effects.

**Study Limitations**

HiPSC-CMs from 2 healthy donors and 1 SQTS patient were used for this study. Differences among individuals cannot be ruled out. The results from the 3 individuals of this study should, from a statistical point of view, not be interpreted as that from the whole population of SQT patients and healthy people. Statistical analyses did not take into account possible correlation between measurements taken from the same individual. hiPSC-CMs possess similarities but also distinct differences in their physiological properties when compared
with adult human cardiomyocytes. In particular, the immature characteristics of iPSC-CMs are a well-known limitation. In addition, in the single-cell studies, many factors such as cell-to-cell interaction, nerve and hormone regulations are not involved. The differences between a single-cell and whole organ system should be considered in interpretation of the experimental data from hiPSC-CMs.

It is known that both Cs+ and the mutation (N588K) reduced the sensitivity of the channel to E-4031. Therefore, the analysis of E-4031-sensitive currents may lead to underestimation of Ikr, especially in SQT-hiPSC-CMs. Furthermore, how Cs+ influences the sensitivity of the mutant HERG channel to E-4031 and how the mutation influences the effects of Cs+ on the sensitivity of the channel to E-4031 still need to be investigated.

Acknowledgments
We gratefully thank C. Liebetrau, M. Grohe, L. Rogge, and Y. Wiegrae for excellent technical assistance. We acknowledge the financial support of the Deutsche Forschungsgemeinschaft and Ruprecht-Karls-Universität Heidelberg within the funding program Open Access Publishing. We thank the China Scholarship Council (CSC) for the financial support for Zhihan Zhao and Xin Li.

Sources of Funding
This study was supported by the DZHK (German Center for Cardiovascular Research), the BMBF (German Ministry of Education and Research) (grants to Borggrefe, Utikal, Wieland, and Zimmermann).

Figure 9. Carbachol increased and quinidine abolished arrhythmic events in SQTS (short QT syndrome)-cardiomyocytes. A and B, Representative Ca2+ transients in a donor (D1) cell in absence (A) and presence (B) of 10 μmol/L carbachol (CCh). C, Percentages of cells showing CCh-induced arrhythmia events. D and E, Representative Ca2+ transients in a SQTS-cell in absence (D) and presence (E) of 10 μmol/L CCh. The red arrows indicate the arrhythmia events. F, Representative Ca2+ transients in a SQTS-cell in presence of 10 μmol/L CCh plus 10 μmol/L quinidine.
Disclosures

None.

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SUPPLEMENTAL MATERIAL
Material and Methods

Ethics statement
The skin biopsies from two healthy donors and one SQTS patient were obtained with written informed consent. The study was approved by the Ethics Committee of Medical Faculty Mannheim, Heidelberg University (approval number: 2009-350N-MA) and by the Ethics Committee of University Medical Center Göttingen (approval number: 10/9/15), and carried out in accordance with the approved guidelines and conducted in accordance with the Helsinki Declaration of 1975, as revised in 1983.

Generation of human iPS cells (hiPSC)

Somatic cell isolation and primary culture
Human fibroblast cultures were derived from skin punch biopsies of the donors. Skin punch biopsy (3-4 mm) was taken aseptically by a surgeon, placed in DMEM (Thermo Fisher Scientific, #11960044) containing 200 U/ml penicillin and 200 µg/ml streptomycin (Thermo Fisher Scientific, #15140122) and transferred to the lab within 48 h. Biopsies were mechanically cut in pieces of 0.5-1 mm, were placed epidermis upside in the cell culture dish and cultured in human fibroblast medium (HFBM) composed of DMEM (Thermo Fisher Scientific, #11960044) supplemented with 10% Fetal Bovine Serum (Thermo Fisher Scientific, #10270106), 1x MEM Non-Essential Amino Acids Solution (Thermo Fisher Scientific, #11140035), 2 mM L-Glutamine (Thermo Fisher Scientific, #25030024), 50 µM β-mercaptoethanol (Serva Electrophoresis, #28625), 10 ng/ml bFGF (Peprotech, #100-18B), 100 U/ml penicillin and 100 µg/ml streptomycin (Thermo Fisher Scientific, #15140122) at
37°C with 5% CO2 atmosphere. Medium was changed every other day. Reprogramming of fibroblasts was performed before passage 3 (p3).

**Generation of hiPSCs**

HiPSC lines isSTQSa1.7 (GOEi091-A.7, here abbreviated as SQTS), isSQTSa1.8 (GOEi091-A.8) and isSQTSa1.15 (GOEi091-A.15) were generated from fibroblasts in feeder free culture conditions using the integration-free CytoTune-iPS 2.0 Sendai Reprogramming Kit (Thermo Fisher Scientific, #A16517) with the reprogramming factors OCT4, KLF4, SOX2, c-MYC according to manufacturer’s instructions with modifications. In brief, 1.5x10^4 early passage fibroblasts were plated in two wells of a Matrigel-coated 24-well plate in HFBM two days before transduction. Cells were transduced at 40-50% confluence with Sendai virus cocktail (hKOS: hc-Myc: hKlf4) at a MOI of 10:10:6 according to the counted cell number of extra well (typically 2.5x10^4 cells/well) in HFBM. Virus was removed after 24 h and HFBM was changed every other day.

HiPSC lines ipWT1.1 (GOEi014-B.1), ipWT1.3 (GOEi014-B.3, here abbreviated as D2) and ipWT1.6 (GOEi014-B.6) were generated from fibroblasts in feeder free culture conditions using the integration-free episomal 4-in-1 CoMiP reprogramming plasmid (Addgene, #63726) with the reprogramming factors OCT4, KLF4, SOX2, c-MYC and short hairpin RNA against p53, as described previously with modifications. In brief, 5x10^5 early passage fibroblasts were used for electroporation with the Nucleofector 2b Device (Lonza) with program P22 or U23 by using the NHDF Nucleofector Kit (Lonza, #VPD-1001) and 2 µg of the reprogramming plasmid. Transfected cells were plated in one well of a Matrigel-coated 6-well plate in HFBM supplemented with 500 µM sodium butyrate (Sigma-Aldrich, #B5887), 2 µM Thiazovivin (Millipore, #420220), 100 U/ml penicillin and 100 µg/ml streptomycin. Medium was changed every other day with HFBM supplemented with 500 µM sodium butyrate.
At d7 post transduction/transfection, cells were replated in various dilutions in Matrigel-coated 6-well plates in HFBM supplemented with 500 µM sodium butyrate and 2 µM Thiazovivin. From d8, medium was changed to E8 medium (Thermo Fisher Scientific, #A1517001) supplemented with 500 µM sodium butyrate (d8-d11) with daily medium change. Cells were monitored for morphology change and appearance of colonies (typically after 2-3 weeks). Individual colonies with iPSC-like morphology were picked mechanically in Matrigel-coated 12-well plates in E8 medium supplemented with 2 µM Thiazovivin. Newly established iPSC lines were passaged with Versene solution (Thermo Fisher Scientific, #15040066) and cultured in E8 medium supplemented with 2 µM Thiazovivin on the first day after passaging in Matrigel-coated plates for at least ten passages before being used for pluripotency characterization and differentiation experiments.

**Spontaneous in vitro differentiation of hiPSCs**

For embryoid body (EB) formation, 5x10⁴ hiPSCs together with 2.5x10⁴ mouse embryonic fibroblasts were plated in each well of a 96-well U-bottom plate in hES medium composed of DMEM-F12 (Thermo Fisher Scientific, #31331028), 15% Knockout Serum Replacement (Thermo Fisher Scientific, #10828028), 1x MEM Non-Essential Amino Acids Solution (Thermo Fisher Scientific, #11140035), 50 µM β-mercaptoethanol (Serva Electrophoresis, #28625) and 2 µM Thiazovivin, the plate was centrifuged at 250 g for 5 min and co-cultures were cultivated in suspension to form multicellular EB aggregates. At d2, medium was changed to differentiation medium composed of IMDM Glutamax (Thermo Fisher Scientific, #31980022), 20% Fetal Bovine Serum (Thermo Fisher Scientific, #10270106), 1x MEM Non-Essential Amino Acids Solution and 450 µM 1-Thioglycerol (Sigma-Aldrich, #M6145) for further 6 days with medium change every other day. At d8, EBs were plated onto 0.1% gelatin-coated 6-well plates and cultured for up to one month in differentiation medium with medium change every other day.
Alkaline phosphatase staining

Alkaline phosphatase activity in hiPSC lines was detected using the Alkaline Phosphatase Kit (Sigma-Aldrich, #86R-1KT) according to the manufacturer’s instructions.

Generation of hiPSC-CMs

Frozen aliquots of hiPSCs were thawed and cultured without feeder cells and differentiated into hiPSC-CMs as described with some modifications. Culture dishes and wells were coated with Matrigel (Corning). Culture medium of hiPSCs was TeSR-E8 (Stemcell Technologies) and for hiPSC-CMs we used RPMI 1640 Glutamax (Life Technologies) containing sodium pyruvate, Penicillin / Streptomycin, B27 (Life Technologies) and ascorbic acid (Sigma Aldrich). Adding of CHIR99021 (Stemgent), BMP-4 (R&D Systems), Activin A (R&D Systems), FGF-2 (MiltenyiBiotec) and IWP-4 (Stemgent) at different time points induced the cells to differentiate into hiPSC-CMs during 3 weeks. During the third week a lactate (Sigma Aldrich) containing RPMI-medium without glucose and glutamine (WKS, Germany) was added to select for cardiomyocyte cells. At 40 to 50 days of culture with basic culture medium, cardiomyocytes were dissociated from 24 well plates and plated on Matrigel-coated 3.5 cm petri dishes for patch-clamp measurements: The cells were incubated with 300µl (150 U) collagenase CLS I (Worthington, Germany) for 40 min at 37°C, washed with PBS and incubated with 0.05% Trypsin-EDTA (Life Technologies) for 4 min at 37°C. After adding of RPMI medium containing 10% FCS, cells were centrifuged at 250 x g for 2 min at room temperature, the supernatant discarded and the cells resuspended with basic culture medium. The cells were plated on the 3.5 cm petri dishes at a density of 2 – 4 x 10⁴ cells/dish for subsequent patch-clamp experiments. A minimum of 3 differentiation experiments of all 3 different iPSC lines were used for each analysis.

Reverse transcriptase-polymerase chain reaction
To quantify the steady-state mRNA expression of the hiPSC-CMs, RNA was reverse transcribed and qPCR was performed as described \(^3\). Gene symbols, RefSeq No. and Cat. No. of the primers used for qPCR analyses in hiPSC-CMs characterization were listed in supplementary table 1 (Table S1). For evaluation of the characteristics of the used hiPSC lines, RT-PCR was performed as follows: Total RNA of hiPSC cultures was isolated using the SV Total RNA Isolation System (Promega, #Z3105) according to manufacturer’s instructions. 100 ng RNA was used for the first-strand cDNA synthesis by using MULV Reverse Transcriptase (Thermo Fisher Scientific, #N8080018) and Oligo d(T)16 (Thermo Fisher Scientific, #N8080128). One-tenth of cDNA was used as PCR template and amplified using the GoTaq G2 DNA polymerase (Promega, #M7845) according to manufacturer’s instructions. The relative expression level of the target gene compared with that of the housekeeping gene, GAPDH, was calculated by the \(2^{-\Delta\Delta CT}\) method. Primer sequences, annealing temperatures and cycles used for RT-PCR analyses of the hiPSC lines were listed in supplementary table 2 (Table S2).

**Immunocytochemical staining**

HiPSC cultures were fixed with Roti-Histofix 4% (Carl Roth, #P087) at RT for 20 min and blocked with 1% bovine serum albumin (BSA; Sigma Aldrich, #F7524) in PBS at 4°C overnight. Primary antibodies were applied in 1% BSA for 1 h at 37°C or overnight at 4°C. Secondary antibodies with minimal cross reactivity were administered in 1% BSA for 1 h at RT. For nuclear or cytosolic proteins (OCT4, SOX2, NANOG, LIN28), cells were permeabilized with 0.1% Triton-X100 (Carl Roth, #3051) in staining solution. Nuclei were stained with 4.8 μM DAPI (Thermo Fisher Scientific, #D1306) for 10 min at RT. Samples were mounted in Fluoromount-G (Thermo Fisher Scientific, #00-4958-02). Antibodies and dilutions used were listed in supplementary table 3 (Table S3). Images were collected using the Axio
Observer Z1 microscopy system (Carl Zeiss) with Axiocam MRm, Plan-Apochromat 20x/0.8 objective or A-Plan 10x/0.25 objective and Axiovision software.

Immunofluorescence staining for KCNH2 channel analysis was performed using appropriate primary antibodies and Alexa Fluor conjugated secondary antibodies (ThermoFisher) according to the manufacturer’s instructions. The primary antibodies used in hiPSC-CMs were a-actinin (Sigma Aldrich), TNNT (Sigma Aldrich) and KCNH2 (Santa Cruz; Alomone). All the other antibodies used for characterization of hiPSC lines are listed in supplementary table 3 (Table S3). Fluorescence measurement was performed according to method described previously using ImageJ 4. Briefly, the cell of interest was selected, and measurement parameters were set to get the area, integrated density and mean gray value. Then a region next to the cell that has no fluorescence was selected as the background. This step was repeated for other cells in the field of view that should be measured. The corrected total cell fluorescence was calculated as (CTCF)= Integrated Density – (Area of selected cell x Mean fluorescence of background readings). Mean fluorescence density was obtained by CTCF/ Area. Pictures were taken using the same exposure parameters. More than 60 cells from 5-8 pictures were selected for analysis. Given that the cells were not permeated during staining and the anti-Kv11.1 (hERG) (extracellular) antibody binding to the extracellular peptide between S1 and S2 domains was used, the staining was considered to come from surface membrane hERG channels.

**Flow cytometry**

HiPSC cultures between passage 10-15 were dissociated with Versene solution into single cells, fixed with Roti-Histofix 4% (Carl Roth, #P087) at RT for 20 min and blocked with 1% bovine serum albumin (BSA; Sigma Aldrich, #F7524) in PBS at 4°C for at least 2 h. Cells were permeabilized with 0.1% Triton-X100 in staining solution and co-incubated with fluorescence-conjugated antibodies against OCT4 and TRA-1-60 at RT for 1 h. Nuclei were co-stained with 8.1 µM Hoechst 33342 (Thermo Fisher Scientific, # H1399). Subsequently, cells were analyzed
using the LSRII flow cytometer (BD Biosciences) and BD FACSDiva software. Gating of cells was applied based on forward scatter area (FSC-A) and sideward scatter area (SSC-A) as well as on gating of single cells (based on DNA signal width). At least 10,000 events were analyzed per sample.

**Measurement of intracellular calcium concentration**

To measure the intracellular Ca\(^{2+}\) concentration (\([\text{Ca}^{2+}]_i\)), cells were loaded with the fluorescent Ca\(^{2+}\)-indicator Fluo-3 AM. First, 1.5 ml PSS (see below) was added into a petri dish with hiPSC-CMs cultured for 2 to 4 days. The following steps were executed under consideration of the light sensitivity of the fluorescent Ca\(^{2+}\) Indicator Fluo-3. Then, 50 µg of the membrane permeable acetoxymethyl ester derivative of Fluo-3 was dissolved in 44 µl of the Pluronic F-127 stock solution (20% w/v in DMSO) to get a 1 mM Fluo-3 AM stock solution, which can be stored at -20 °C for a maximum of 1 week. Next, 15 µl of the Fluo-3 AM stock solution were added into 1.5 ml PSS resulting in a final concentration of 10 µM Fluo-3 and the dish was agitated carefully. The cells were incubated at room temperature for 10 minutes in an optically opaque box to protect from light. Thereafter, the PSS was carefully sucked out and discarded and the cells were washed with PSS for 4-5 times. Finally, the cells in PSS were kept at room temperature for about 30 minutes for de-esterification before measurements. After de-esterification the fluorescence of the cells was measured by using Cairn Optoscan calcium imaging system (Cairn Research, UK). Fluorescence is excited by 488 nm and emitted at 520 nm. Changes in \([\text{Ca}^{2+}]_i\) were described by

\[
[\text{Ca}^{2+}]_i = k_d \left( \frac{F}{F_{\text{max}} - F} \right)
\]

where \(k_d=\)dissociation constant of Fluo-3 (400-nmol/L), \(F=\)Fluo-3 fluorescence; \(F_{\text{max}}=\)Ca\(^{2+}\)-saturated fluorescence obtained at the end of each experiment by damaging the cell with the patch pipette and measuring the increase of fluorescence to maximum \(^5\).
**Patch-clamp**

Standard patch-clamp recording techniques were used to measure the $I_{\text{Na}}$, $I_{\text{CaL}}$, $I_{\text{to}}$, $I_{\text{Kr}}$, $I_{\text{Ks}}$, $I_{\text{K1}}$ and action potential (AP) in the whole-cell configuration at room temperature. Patch electrodes were pulled from borosilicate glass capillaries (MTW 150F; world Precision Instruments, Inc., Sarasota, FL) using a DMZ-Universal Puller (Zeitz-InstrumenteVertriebs GmbH, Martinsried, Germany) and filled with pre-filtered pipette solution (see below). Pipette resistance ranged from 1-2 MΩ. Electrode offset potentials were zero-adjusted before a Giga-seal was formed. After a Giga-seal was obtained, fast capacitance was first compensated and then the membrane under the pipette tip was disrupted by negative pressure to establish the whole-cell configuration. To determine the cell capacitance, a voltage pulse from -80 to -85 was given to record the cell capacitance transient current. Then the area under transient current curve was integrated and divided by 5 mV to get the whole cell capacitance in pF. Thereafter the membrane capacitance ($C_m$) and series resistance ($R_s$) were compensated (60-80%). Liquid junction potentials were not corrected. Signals were acquired at 10 kHz and filtered at 2 kHz with the Axon 200B amplifier and Digidata 1440A digitizer hardware as well as pClamp10.2 software (Molecular Devices, Sunnyvale, CA). Myocytes were held at -80 mV for $I_{\text{Na}}$, $I_{\text{to}}$, $I_{\text{CaL}}$ and at -40 mV for $I_{\text{Kr}}$, $I_{\text{Ks}}$, and $I_{\text{K1}}$ measurements. For $I_{\text{Na}}$ measurements, currents were elicited by using depolarizing pulses of 400 ms from -100 mV to +50 mV with 10 mV increments at 0.5 Hz. For measurements of $I_{\text{CaL}}$, cells were depolarized from -80 mV to -40 mV for 500 ms and then to different potentials (-100 to +60 mV) for 500 ms. For $I_{\text{Kr}}$ recordings, cells were depolarized from -20 to +80 mV for 2s and repolarized to -30 mV. For $I_{\text{Ks}}$ recordings, cells were depolarized from -40 mV to +50 mV for 4s and repolarized to -40 mV. For $I_{\text{to}}$ recordings, currents were evoked by pulses from -80 mV to +80 mV for 500 ms. For $I_{\text{K1}}$, currents were elicited by pulses from -120 mV to +60 mV for 500 ms. Measured currents were normalized to the membrane capacitance. Current-voltage (I-V) relationships were generated by plotting the current density against voltages. The TTX sensitive late $I_{\text{Na}}$ was measured as the area under
the current curve integrated from 50 to 350 ms after the beginning of the depolarization pulse. E-4031 (3 µM) was used for isolating \( I_{Kr} \), chromanol 293 B (10 µM) for \( I_{Ks} \), 4-AP (5 mM) for \( I_{to} \), BaCl\(_2\) (100 µM) for \( I_{K1} \). The blocker-sensitive currents were used for comparison between donor- and sQT-cells. To minimize the effects of run-down of recorded currents on the results of experiments, we carefully monitored the time-dependent change of currents. Recordings were started after the current reached a steady state, normally within 3 to 5 minutes.

The bath solution for late sodium current contained (mmol/l): 135 NaCl, 20 CsCl, 1.8 CaCl\(_2\), 1 MgCl\(_2\), 10 Hepes, 10 glucose, 0.001 nifedipine, pH 7.4 (CsOH). Microelectrodes were filled with (mmol/l): 10 NaCl, 135 CsCl, 2 CaCl\(_2\), 3 MgATP, 2 TEA-Cl, 5 EGTA and 10 HEPES (pH7.2 CsOH).

The bath solution for \( I_{CaL} \) recordings contained (mmol/l): 140 TEA-Cl, 5 CaCl\(_2\), 1 MgCl\(_2\), 10 Hepes, 0.01 TTX, 2 4-AP, pH 7.4 (CsOH). Microelectrodes were filled with (mmol/l): 10 NaCl, 135 CsCl, 2 CaCl\(_2\), 3 MgATP, 2 TEA-Cl, 5 EGTA and 10 HEPES (pH7.2 CsOH).

The bath solution (PSS) for \( I_{to} \), \( I_{Ks} \), \( I_{K1} \) and AP measurements contained (mmol/l): 130 NaCl, 5.9 KCl, 2.4 CaCl\(_2\), 1.2 MgCl\(_2\), 11 glucose, 10 HEPES, pH 7.4 (NaOH). For the \( I_{to} \) measurements, 10 µM nifedipine, 10 µM TTX and 1 µM E-4031 were added in the bath solution to block \( I_{CaL} \), \( I_{Na} \) and \( I_{Kr} \). For \( I_{Ks} \) measurements, 2 mM 4-AP and 1 µM E-4031 were added. For \( I_{K1} \) measurements the concentration of KCl was increased to 20 mM. The pipette solution contains 10 mM HEPES, 126 mM KCl, 6 mM NaCl, 1.2 mM MgCl\(_2\), 5 mM EGTA, 11 mM glucose and 1 mM MgATP, pH 7.4 (KOH).

To improve \( I_{Kr} \) measurements, the Cs\(^{2+}\) currents conducted by the KCNH2 channels were measured. External solution for Cs\(^{2+}\) currents (mmol/L): 140 CsCl, 2 MgCl\(_2\), 10 HEPES, 10 Glucose, pH=7.4 (CsOH). Pipette solution: 140 CsCl, 2 MgCl\(_2\), 10 HEPES, 10 EGTA, pH=7.2 (CsOH).
**Statistics**

If not otherwise indicated data are shown as mean ± SEM and were analyzed using InStat© (GraphPad, San Diego, USA) and SigmaPlot 11.0 (Systat GmbH, Germany). By analyzing the data with the Kolmogorov Smirnov test it was decided whether parametric or non-parametric tests were used for analysis. Student’s t-test and the Mann–Whitney U-test were used to compare continuous variables with normal and non-normal distributions, respectively. The chi-squared-test for independence was used to compare categorical variables. For parametric data one-way ANOVA with Bonferroni post-test for multiple comparisons was performed. For non-parametric data the Kruskal-Wallis test with Dunn’s multiple comparisons post-test was used. Paired t-test was used for comparisons of data before and after application of a drug. $p<0.05$ (two-tailed) was considered significant.
Table S1. Gene symbols, RefSeq No. and Cat. No. of the primers used for qPCR analyses in hiPSC-CMs characterization.

| Gene symbol | RefSeq No.   | Cat. No. Primers |
|-------------|--------------|------------------|
| CACNA1C (L-type Ca2+ channel) | NM_000719 | PPH01378G |
| KCND3 (Ito, Kv4.3) | NM_004980 | PPH06923A |
| KCNH2 (IKr, Kv11.1) | NM_000238 | PPH01660A |
| KCNJ2 | NM_000891 | PPH01618E |
| KCNQ1 (Iks, Kv7.1) | NM_000218 | PPH01419A |
| SCN5A (Na+ channel, Nav1.5) | NM_000335 | PPH01671F |

RefSeq No. : GenBank NCBI Reference Sequences  
Cat. No. Primers: Qiagen RT² qPCR Primer Assays
Table S2. Primer sequences, annealing temperatures and cycles used for RT-PCR analyses in hiPSC pluripotency characterization.

| Primer | Primer for          | Primer rev               | Length bps | Temp /Cycles |
|--------|---------------------|--------------------------|------------|--------------|
| OCT4   | GACAACAATGAAAATCTTCAGAGA | TTCTGGCGCCGGTTTACAGAACCA | 218        | 58°C /36 cycles |
| SOX2   | ATGCACCGCTACGACGTGA  | CTTTGACACCCTCCCATT      | 437        | 58°C /30 cycles |
| NANO   | AGTCCCAAAGGCAACAACCACTTC | ATCTGCTGGAGGCTAGGTTATTCCTTC | 164        | 64°C /36 cycles |
| LIN28  | AGTAAGCTGCACATGGAAGG | ATTGTGGCTCAATCTGTGC      | 410        | 58°C /36 cycles |
| FOXD3  | GTGAAGCCGCTTTAATCCTAC | CCAGAAGCTCTGCATCATGAG    | 353        | 58°C /38 cycles |
| GDF3   | TTCGCTCTCCAGACCAAGGTTC | TACCACAGCAGTGAAGATGAAAAACAC | 311        | 58°C /32 cycles |
| AFP    | ACTCACTAAACCTGGTGTG | GAAATCTGCAATGACAGCCTCA   | 255        | 58°C /30 cycles |
| ALB    | CTTTGACACATGAAGTTGGAACCAAC | CAGCAGTACACCCATTTACCATAGG | 355        | 62°C /35 cycles |
| TNNT2  | GACAGAGCGGAAATGTAAGGA | GTAAGAGGCCAGGCTTAT      | 305        | 58°C /30 cycles |
| MYH6   | GTGATTGCTGAAACCGAGTGAA | GCAAGTACCTGAGAAGATGACACGCT | 413         | 58°C /38 cycles |
| TH     | GCCGCTCATTGGGCGCAGG | CAAACACCTTCACAGCTCG     | 215        | 58°C /34 cycles |
| MAP2   | CCACCTGAGATAAGGATCA  | GGCTTACTTTGCTCTCTGA     | 482        | 55°C /32 cycles |
| GAPDH  | AGAGGCAGGGATGATGTTCT | TCTGCTGATGCCCCATGTT     | 258        | 58°C /30 cycles |
**Table S3.** Antibodies and dilutions used for immunocytochemistry of iPSC pluripotency characterization.

| Primary antibody | Type                  | Dilution | Supplier                        |
|------------------|-----------------------|----------|---------------------------------|
| hOCT3/4          | polyclonal goat IgG   | 1:40     | R&D Systems, #AF1759            |
| hSOX2            | monoclonal mouse IgG1 | 1:200    | Thermo Fisher Scientific, #MA1-014 |
| hNANOG           | polyclonal rabbit IgG | 1:100    | Thermo Fisher Scientific, #PA1-097 |
| hLIN28           | polyclonal goat IgG   | 1:300    | R&D Systems, #AF3757            |
| hSSEA4           | monoclonal mouse IgG3 | 1:100    | Thermo Fisher Scientific, #MA1-021 |
| hTRA-1-60        | monoclonal mouse IgM  | 1:200    | Abcam, #ab16288                 |
| hAFP             | polyclonal rabbit IgG | 1:100    | DAKO, #A0008                    |
| α-SMA            | monoclonal mouse IgG2a| 1:3000   | Sigma-Aldrich, #A2547           |
| β-III-Tubulin    | monoclonal mouse IgG2A| 1:2000   | Covance, #MMS-435P              |

| Secondary antibody | Type                  | Dilution | Supplier                        |
|--------------------|-----------------------|----------|---------------------------------|
| Alexa Fluor 488    | polyclonal donkey α-mouse IgG | 1:1000   | Thermo Fisher Scientific, #A21202  |
| Alexa Fluor 555    | polyclonal donkey α-goat IgG  | 1:1000   | Thermo Fisher Scientific, #A21432  |
| Alexa Fluor 555    | polyclonal donkey α-mouse IgG  | 1:1000   | Thermo Fisher Scientific, #A31570  |
| Alexa Fluor 555    | polyclonal donkey α-rabbit IgG | 1:1000   | Thermo Fisher Scientific, #A31572  |
| FITC               | polyclonal goat α-mouse IgM  | 1:200    | Jackson Immuno, #115-097-020     |

**Flow Cytometry**

|                        | Dilution | Supplier |
|------------------------|----------|----------|
| Alexa Fluor 488 mouse anti-human TRA-1-60 | 1:50    | BD Biosciences, #560173 |
| Alexa Fluor 647 mouse anti-OCT3/4     | 1:50    | BD Biosciences, #560329 |
Table S4. Action potential parameters.

| group          | RP (mV)    | APA (mV)  | Vmax (V/s) | APD50 (ms) | APD90 (ms) | n  |
|----------------|------------|-----------|------------|------------|------------|----|
| D1             | -79.07±1.17| 135.60±2.63| 32.29±2.67 | 148.07±14.19| 320.17±19.08| 28 |
| D2             | -80.17±0.73| 126.66±3.08| 28.52±2.34 | 149.34±9.68 | 289.21±13.77| 29 |
| SQTS           | -78.07±0.76| 127.31±2.74| 30.86±2.28 | *93.86±5.15 | *187.90±6.91| 29 |
| SQTS (Ctr)     | -79.92±0.51| 131.00±4.47| 32.83±3.97 | 91.75±10.57 | 203.17±15.76| 12 |
| Quinidine      | -80.42±0.70| 127.00±3.63| #28.17±3.22| #130.83±16.49| #324.17±42.69| 12 |
| SQTS (Ctr)     | -81.58±0.67| 122.25±4.39| 26.00±3.58 | 94.92±15.30 | 204.83±31.11| 12 |
| Metoprolol     | -81.83±0.69| 123.33±4.18| 25.17±3.24 | 91.08±13.14 | 218.33±36.14| 12 |
| SQTS (Ctr)     | -79.09±0.79| 133.36±4.96| 35.45±3.94 | 99.27±9.88 | 219.36±16.45| 11 |
| Sotalol        | -78.64±0.66| 133.18±4.68| 35.18±3.81 | 93.73±9.38 | 230.45±18.25| 11 |

*p<0.05 versus D1; #p<0.05 versus Ctr
Figure S1. The generated hiPSC lines from the patient with short QT syndrome (SQTS) display pluripotent characteristics.

(A) The hiPSC (human induced pluripotent stem cell) lines isSQTSa1.7, isSQTSa1.8 and isSQTSa1.15 generated from skin fibroblasts of the SQTS-patient display a typical morphology for human pluripotent stem cells (upper panel) and are positive for alkaline phosphatase (lower panel). (B) In comparison to fibroblasts, generated hiPSC lines show expression of endogenous pluripotency markers SOX2 (sex determining region Y-box 2), OCT4 (octamer-binding transcription factor 4), NANOG (pron. nanOg, homeobox protein), LIN28 (lin-28 homolog A), FOXD3 (Forkhead Box D3) and GDF3 (Growth differentiation factor-3) at mRNA level proven
by RT-PCR (reverse transcription-polymerase chain reaction). Human embryonic stem cells (hESCs) were used as positive control, mouse embryonic fibroblasts (MEFs) were used as negative control. (C) Generated iPSC lines express pluripotency markers OCT4, SOX2, NANOG, LIN28, SSEA4 (stage-specific embryonic antigen 4) and TRA-1-60 as shown by immunofluorescence staining. Nuclei are co-stained with DAPI (4',6-diamidino-2-phenylindole). (D) Spontaneous differentiation potential of generated iPSC lines was analysed by embryoid body (EB) formation. Germ layer-specific genes like α-fetoprotein (AFP) and albumin (ALB) (endoderm), cTNT (cardiac troponin T) and α-MHC (mesoderm), and tyrosine hydroxylase (TH) and MAP2 (ectoderm) are expressed in a developmentally controlled manner during differentiation of EBs (days 0, 8, or 8+25), whereas endogenous OCT4 expression is decreased during spontaneous differentiation. MEFs were used as negative control. (E) Immunocytochemical staining of spontaneously differentiated hiPSC lines shows expression of endodermal marker AFP, mesodermal-specific α-SMA (α-smooth muscle actin) and ectodermal βIII-tubulin. Nuclei are co-stained with DAPI. Scale bars: 100 µm.
Figure S2. The generated hiPSC lines from a healthy donor display pluripotent characteristics.

(A) The hiPSC (human induced pluripotent stem cell) lines ipWT1.1, ipWT1.3 and ipWT1.6 generated from skin fibroblasts of a healthy donor (D2) display a typical morphology for human pluripotent stem cells (upper panel) and are positive for alkaline phosphatase (lower panel). (B) In comparison to fibroblasts, generated hiPSC lines show expression of endogenous pluripotency markers SOX2 (sex determining region Y-box 2), OCT4 (octamer-binding transcription factor 4), NANOG (pron. nanOg, homeobox protein), LIN28 (lin-28 homolog A),
FOXD3 (Forkhead Box D3) and GDF3 (Growth differentiation factor-3) at mRNA level proven by RT-PCR (reverse transcription-polymerase chain reaction). Human embryonic stem cells (hESCs) were used as positive control, mouse embryonic fibroblasts (MEFs) were used as negative control. (C) Generated hiPSC lines express pluripotency markers OCT4, SOX2, NANOG, LIN28, SSEA4 (stage-specific embryonic antigen 4) and TRA-1-60 as shown by immunofluorescence staining. Nuclei are co-stained with DAPI (4',6-diamidino-2-phenylindole). (D) Spontaneous differentiation potential of generated hiPSC lines was analysed by embryoid body (EB) formation. Germ layer-specific genes like α-fetoprotein (AFP) and albumin (ALB) (endoderm), cTNT (cardiac troponin T) and α-MHC (mesoderm), and tyrosine hydroxylase (TH) and MAP2 (ectoderm) are expressed in a developmentally controlled manner during differentiation of EBs (days 0, 8, or 8+25), whereas endogenous OCT4 expression is decreased during spontaneous differentiation. MEFs were used as negative control. (E) Immunocytochemical staining of spontaneously differentiated hiPSC lines shows expression of endodermal marker AFP, mesodermal-specific α-SMA (α-smooth muscle actin) and ectodermal βIII-tubulin. Nuclei are co-stained with DAPI. Scale bars: 100 μm.
Figure S3. Slowly activating delayed rectifier currents (I\(_{Ks}\)) in donor-and SQTS (short QT syndrome)-cells.

I\(_{Ks}\) (slowly activating delayed rectifier potassium channel) were evoked by the indicated protocol (B) in absence (control) and presence of a channel blocker. Chromanol 293B (10\(\mu\)M) was used to isolate I\(_{Ks}\) from other currents. (A) Representative I\(_{Ks}\) at +40 mV recorded in a cell from a donor (D2) and the patient (SQTS) with and without (Ctr) 293B. (B) 293B-sensitive currents. (C) I-V curves of I\(_{Ks}\) from the donor- and SQTS-cells. n, number of cells. *p<0.05 versus donor cells.
Figure S4. The transient outward currents ($I_{to}$) and inward rectifier currents ($I_{K1}$) in donor- and SQTS (short QT syndrome)-cells.

$I_{to}$ and $I_{K1}$ were evoked by the indicated protocol (A and C) in absence (Ctr) and presence of a channel blocker. 4-aminopyridine (4-AP, 5mM) and BaCl$_2$ (0.1 mM) was used to isolate $I_{to}$ and $I_{K1}$ from other currents. (A) Representative $I_{to}$ at +40 mV recorded in a cell from the donors (D1 and D2) and the patient (SQTS). (B) I-V curves of $I_{to}$ from the donor- and SQTS-cells. (C) Representative $I_{K1}$ at -120 mV recorded in a cell from the donors (D1 and D2) and the patient (SQTS). (D) I-V curves of $I_{K1}$ from the donor- and SQTS-cells. n, number of cells.
Figure S5. The L-type calcium channel currents ($I_{CaL}$) in donor-and SQTS (short QT syndrome)-cells.

$I_{CaL}$ was evoked by the indicated protocol (A). (A) Representative $I_{CaL}$ at different potentials recorded in a cell from the donors (D1 and D2) and the patient (SQTS). (B) I-V (current-voltage relationship) curves of $I_{CaL}$ from the donor- and SQTS-cells.
Figure S6. Ikr recorded in hiPSC (short QT syndrome)-CMs with a holding potential of -80 mV.

Ikr (rapidly activating delayed rectifier potassium channel) was evoked by 2 s depolarizing pulses from -70 to +70 mV with 10 mV increments and repolarized to the holding potential of -80 mV. (A) Representative traces of I_{Kr} in a donor (D2) cell. (B) Representative traces of I_{Kr} in a SQTS (SQT) cell.
APs (action potentials) were recorded in a cell before (control) and after application of a drug. (A)-(C) Mean values of maximal depolarization speed (Vmax), resting potential (RP) and amplitude of AP (APA) in absence and presence of quinidine. (D)-(F) Mean values of maximal depolarization speed (Vmax), resting potential (RP) and amplitude of AP (APA) in absence and presence of sotalol. (G)-(I) Mean values of maximal depolarization speed (Vmax), resting potential (RP) and amplitude of AP (APA) in absence and presence of metoprolol. n, number of cells.
Figure S8. Effects of antiarrhythmic drugs on action potential duration (APD) in SQTS (short QT syndrome)-cardiomyocytes.

(A) Mean values of APD at 50% repolarization (APD50) in absence (control) and presence of 10 µM quinidine. (B) Mean values of APD at 90% repolarization (APD90) in absence (control) and presence of 10 µM quinidine. (C) Representative traces of action potentials in absence (control) and presence of 10 µM quinidine. (D) Mean values of APD at 50% repolarization (APD50) in absence (control) and presence of 100 µM sotalol. (E) Mean values of APD at 90% repolarization (APD90) in absence (control) and presence of 100 µM sotalol. (F) Representative traces of action potentials in absence (control) and presence of 100 µM sotalol. (G) Mean values of APD at 50% repolarization (APD50) in absence (control) and presence of 10 µM metoprolol. (H) Mean values of APD at 90% repolarization (APD90) in absence (control) and presence of 10 µM metoprolol. (I) Representative traces of action potentials in absence (control) and presence of 10 µM metoprolol. Values given are mean ± SEM (standard error of the mean). n, number of cells. ns, p>0.05.
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