

Recessive cardiac phenotypes in induced pluripotent stem cell models of Jervell and Lange-Nielsen syndrome: Disease mechanisms and pharmacological rescue

Miao Zhang^{a,b,1}, Cristina D'Aniello^{c,1}, Arie O. Verkerk^d, Eva Wrobel^e, Stefan Frank^{a,b}, Dorien Ward-van Oostwaard^f, Ilaria Piccini^e, Christian Freund^c, Jyoti Rao^{a,b}, Guiscard Seebohm^e, Douwe E. Atsma^f, Eric Schulze-Bahr^e, Christine L. Mummery^c, Boris Greber^{a,b,2}, and Milena Bellin^{c,2}

^aHuman Stem Cell Pluripotency Group, Max Planck Institute for Molecular Biomedicine, D-48149 Münster, Germany; ^bChemical Genomics Centre of the Max Planck Society, 44227 Dortmund, Germany; Departments of ^cAnatomy and Embryology and ^dCardiology, Leiden University Medical Center, Leiden, 2300 RC, The Netherlands; ^eDepartment of Anatomy, Embryology and Physiology, Heart Failure Research Center, Academic Medical Center, Amsterdam, 1105 AZ, The Netherlands; and ^fInstitute for Genetics of Heart Diseases, Department of Cardiovascular Medicine, University Hospital Münster, D-48149 Münster, Germany

Edited by R. Michael Roberts, University of Missouri, Columbia, MO, and approved November 7, 2014 (received for review October 15, 2014)

Jervell and Lange-Nielsen syndrome (JLNS) is one of the most severe life-threatening cardiac arrhythmias. Patients display delayed cardiac repolarization, associated high risk of sudden death due to ventricular tachycardia, and congenital bilateral deafness. In contrast to the autosomal dominant forms of long QT syndrome, JLNS is a recessive trait, resulting from homozygous (or compound heterozygous) mutations in *KCNQ1* or *KCNE1*. These genes encode the α and β subunits, respectively, of the ion channel conducting the slow component of the delayed rectifier K^+ current, I_{Ks} . We used complementary approaches, reprogramming patient cells and genetic engineering, to generate human induced pluripotent stem cell (hiPSC) models of JLNS, covering splice site (c.478-2A>T) and missense (c.1781G>A) mutations, the two major classes of JLNS-causing defects in *KCNQ1*. Electrophysiological comparison of hiPSC-derived cardiomyocytes (CMs) from homozygous JLNS, heterozygous, and wild-type lines recapitulated the typical and severe features of JLNS, including pronounced action and field potential prolongation and severe reduction or absence of I_{Ks} . We show that this phenotype had distinct underlying molecular mechanisms in the two sets of cell lines: the previously unidentified c.478-2A>T mutation was amorphic and gave rise to a strictly recessive phenotype in JLNS-CMs, whereas the missense c.1781G>A lesion caused a gene dosage-dependent channel reduction at the cell membrane. Moreover, adrenergic stimulation caused action potential prolongation specifically in JLNS-CMs. Furthermore, sensitivity to proarrhythmic drugs was strongly enhanced in JLNS-CMs but could be pharmacologically corrected. Our data provide mechanistic insight into distinct classes of JLNS-causing mutations and demonstrate the potential of hiPSC-CMs in drug evaluation.

Jervell and Lange-Nielsen syndrome | long QT syndrome | human induced pluripotent stem cells | disease modeling | *KCNQ1*

Jervell and Lange-Nielsen syndrome (JLNS) is a rare, autosomal recessive disease characterized by congenital bilateral deafness, severe QT interval prolongation on the electrocardiogram (ECG), polymorphic ventricular arrhythmias, syncope, and high risk of sudden death (1, 2). JLNS results from homozygous (or compound heterozygous) mutations in the *KCNQ1* or *KCNE1* genes. These encode the α - and β -subunits, respectively, of the ion channel conducting the slow component of the delayed rectifier K^+ current (I_{Ks}) (3, 4). Another long QT condition termed Romano-Ward syndrome (RWS) is, by contrast, an autosomal-dominant form of QT interval prolongation without deafness, caused by heterozygous mutations in 16 different genes, including *KCNQ1* (*LQT1*) and *KCNE1* (*LQT5*) (5–7). However, the recessive JLNS is among the most severe forms of the disease, together with Timothy syndrome and a long QT syndrome variant caused by calmodulin mutations (8, 9). JLNS patients usually have severe clinical symptoms, early disease onset (~12 mo old), and require

aggressive interventions because of the limited efficacy of β -receptor blockers (2).

JLNS patients with *KCNQ1* mutations usually display longer QT intervals and higher risk for arrhythmic events than those with *KCNE1* mutations (2). Attempts to associate the type of mutation (e.g., missense, nonsense, frameshift) in *KCNQ1* with the RWS or JLNS phenotype have proven challenging. In general, however, missense mutations with a dominant-negative effect on the tetrameric *KCNQ1* channel tend to cause RWS, whereas JLNS is frequently caused by nonsense and frameshift mutations (10–13). However, exceptions exist in that missense mutations can also result in JLNS (14). Furthermore, there are rare but well-documented cases of symptoms in heterozygous carriers of JLNS mutations (11, 15–17).

Human induced pluripotent stem cells (hiPSCs) are already proving to provide powerful cellular models to study both genetic and sporadic diseases in humans (18). Several cardiac ion channel diseases have been investigated by using hiPSC-derived cardiomyocytes (hiPSC-CMs), including distinct subtypes of RWS

Significance

There are few laboratory models that recapitulate human cardiac disease. Here, we created human cell models for Jervell and Lange-Nielsen syndrome (JLNS) in vitro, based on human induced pluripotent stem cells (hiPSCs). JLNS is one of the most severe disorders of heart rhythm and can cause sudden death in young patients. JLNS is inherited recessively and is caused by homozygous mutations in the slow component of the delayed rectifier potassium current, I_{Ks} . Cardiomyocytes (CMs) from two independent sets of patient-derived and engineered hiPSCs showed electrophysiological defects that reflect the severity of the condition in patients. Our work allowed better understanding of the mechanisms of recessive inheritance. Furthermore, JLNS-CMs showed increased sensitivity to proarrhythmic drugs, which could be rescued pharmacologically, demonstrating the potential of hiPSC-CMs in drug testing.

Author contributions: E.S.-B., C.L.M., B.G., and M.B. designed research; M.Z., C.D., A.O.V., E.W., S.F., D.W.-v.O., I.P., C.F., J.R., B.G., and M.B. performed research; G.S., D.E.A., and E.S.-B. contributed new reagents/analytic tools; M.Z., C.D., A.O.V., E.W., S.F., I.P., B.G., and M.B. analyzed data; and C.D., A.O.V., E.S.-B., C.L.M., B.G., and M.B. wrote the paper.

Conflict of interest statement: C.L.M. is cofounder and advisor of Pluriomics.

This article is a PNAS Direct Submission.

Freely available online through the PNAS open access option.

¹M.Z. and C.D. contributed equally to this work.

²To whom correspondence may be addressed. Email: boris.greber@mpi-muenster.mpg.de or m.bellin@lumc.nl.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1419553111/-DCSupplemental.

(LQT1, LQT2, LQT3, and LQT8) (19–21). Here, we report and analyze independent hiPSC models for the severe and recessively inherited JLNS. Two JLNS-causing mutations were investigated: the novel *c.478-2A>T* and the previously described *c.1781G>A* single nucleotide exchanges (22). Compared with heterozygous and wild-type (wt) controls, cardiomyocytes (CMs) of both JLNS models showed severe functional abnormalities caused by complete or near-complete loss of I_{Ks} . Although disease phenotypes in the homozygous *c.478-2A>T* and *c.1781G>A* cells were similar, distinct loss-of-function molecular mechanisms (strictly recessive and gene dosage-dependent, respectively) were mediated by the two mutations. JLNS-CMs were also highly sensitive to adrenergic and proarrhythmic stress, which could be exploited in future drug safety pharmacology for identifying high-risk individuals. Conversely, arrhythmia phenotypes could be prevented by pharmacological treatment, highlighting the value of hiPSC-CMs in drug testing.

Results

Generation of hiPSC Lines from Patients with *KCNQ1* Mutations.

Fibroblasts were obtained from patients with different *KCNQ1*

mutations, as follows: (i) a JLNS patient with the homozygous single nucleotide *c.478-2A>T* mutation at the 3' end of intron 2; (ii) an asymptomatic heterozygous carrier from the same family; (iii) a LQT1 patient with the heterozygous missense *c.1781G>A* mutation in exon 15 (Fig. 1A and Fig. S1A–D). Fibroblasts were also derived from two gender-matched healthy individuals (wt1 and wt2; Fig. S1C and D and SI Materials and Methods). The newly described *c.478-2A>T* mutation would possibly lead to a splice defect, resulting in the skipping of exon 3 with concomitant frame shift and appearance of a premature stop codon (E160fs+138X). The *c.1781G>A* *KCNQ1* mutation causes substitution of an arginine with a glutamine residue at position 594 of the coding sequence (R594Q) (Fig. 1B). Sendai viruses carrying the four transcription factors *OCT4*, *SOX2*, *KLF4*, and *MYC* (23) were used to generate hiPSCs. The resulting lines showed typical human embryonic stem cell (hESC) morphology and growth characteristics, with erasure of the Sendai vectors upon passage (Fig. S1E–H). One line per genotype was selected for further experiments. hiPSCs showed activation of endogenous pluripotency markers (Fig. 1C–F), were pluripotent as assessed by

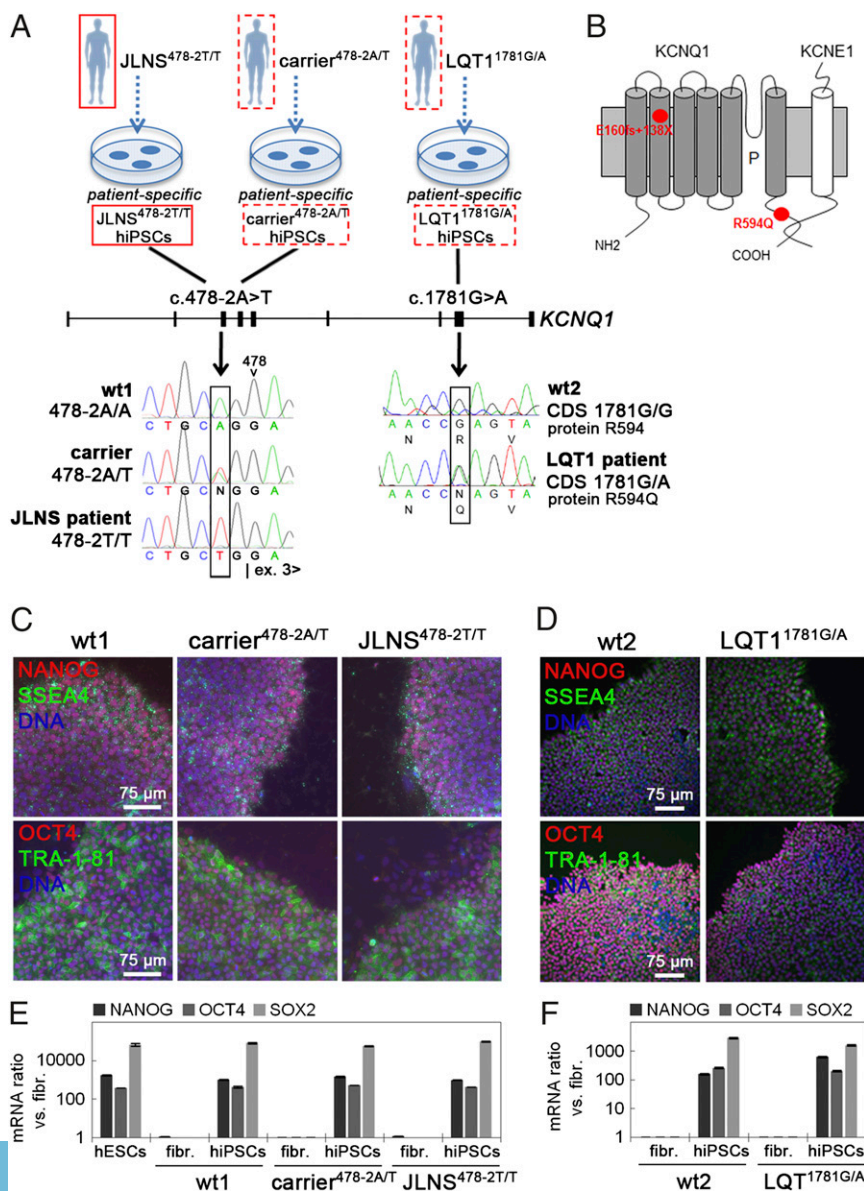


Fig. 1. Generation of hiPSCs from patients with *KCNQ1* mutations. (A) Sequencing of the *KCNQ1* gene identified the *c.478-2A>T* mutation at the splice acceptor site of intron 2 in the JLNS patient and the heterozygous carrier and the heterozygous *c.1781G>A* mutation in exon 15 of the LQT1 patient. The genotypes of the two healthy controls wt1 and wt2 are shown as reference. (B) The *c.478-2A>T* mutation is predicted to result in skipping of exon 3, with a concomitant reading frame shift from position 160 onwards, giving rise to a premature stop codon (E160fs+138X). The *c.1781G>A* mutation results in the substitution of an arginine with a glutamine at position 594 (R594Q) in the C-terminal domain of the *KCNQ1* protein. (C and D) Immunofluorescence analysis of the pluripotency-associated markers NANOG, SSEA4, OCT4, and TRA1-81 in wt1, carrier^{478-2A/T}, and JLNS^{478-2T/T} (C), and in wt2 and LQT1^{1781G/A} hiPSCs (D). (E and F) qPCR expression analysis of the endogenous pluripotency genes *NANOG*, *OCT4*, and *SOX2* in the indicated hiPSC lines.

spontaneous differentiation into derivatives of the three germ layers (Fig. S2 A and B), displayed hESC-like transcriptomes (24) (Fig. S2 C and D), and had normal karyotypes (Fig. S2 E and F). These results indicated successful integration-free reprogramming of all lines.

Generation of Isogenic Pairs of JLNS Human Pluripotent Stem Cells.

To be able to assess the impact of the c.478-2A>T mutation on an independent genetic background, we used the CRISPR/Cas9 system to generate isogenic pairs of wt and homozygous c.478-2A>T hESCs (JLNS^{ts/ts}) by disrupting the intron 2-exon 3 boundary of *KCNQ1* (25) (Fig. 2 A and B). Moreover, we sought to generate a JLNS model from the heterozygous c.1781G>A hiPSCs, by inserting this mutation into the wt allele of the LQT1^{1781G/A} line by using homologous recombination (Fig. 2 C and D). This approach resulted in the isogenic homozygous mutated JLNS^{1781A/A} hiPSC line. Correct targeting in this homozygous clone was confirmed by PCR and DNA sequencing (Fig. S1J). The JLNS^{1781A/A} hiPSCs were further characterized as above, with respect to self-renewal and pluripotency features (Fig. S2 B, D, and F). These cell lines complemented the patient-derived hiPSCs for further investigation.

Differentiation into the Cardiac Lineage. The two independent, validated sets of cell lines (wt1/carrier^{478-2A/T}/JLNS^{478-2T/T} hiPSC and wt2/LQT1^{1781G/A}/JLNS^{1781A/A} hiPSC) were induced to differentiate into CMs by using directed differentiation protocols (SI Materials and Methods). Beating areas were visible from day 6–10 onwards. CMs stained positively for the cardiac sarcomeric proteins MLC2a, MLC2v, troponin I, and α -actinin (Fig. 2 E and F) and expressed genes encoding cardiac ion channels (Fig. 2 G and H). *KCNQ1* is an imprinted gene that is monoallelically expressed during early development, but later, in the heart, expression becomes biallelic (26). Hence, *KCNQ1* imprinting could potentially interfere with proper manifestation of the disease phenotype in hiPSC-CMs. We therefore examined the imprinting status of *KCNQ1* upon cardiac differentiation, by taking advantage of a single-nucleotide polymorphism allowing the two *KCNQ1* alleles in wt1 hiPSCs to be distinguished. Undifferentiated hiPSCs displayed a strictly monoallelic expression pattern, confirming the imprinted status of the gene. However, after differentiation into CMs, expression became biallelic (Fig. 2I). These data suggested that the differentiation methods used here should allow faithful monitoring of differences between heterozygous and homozygous CMs.

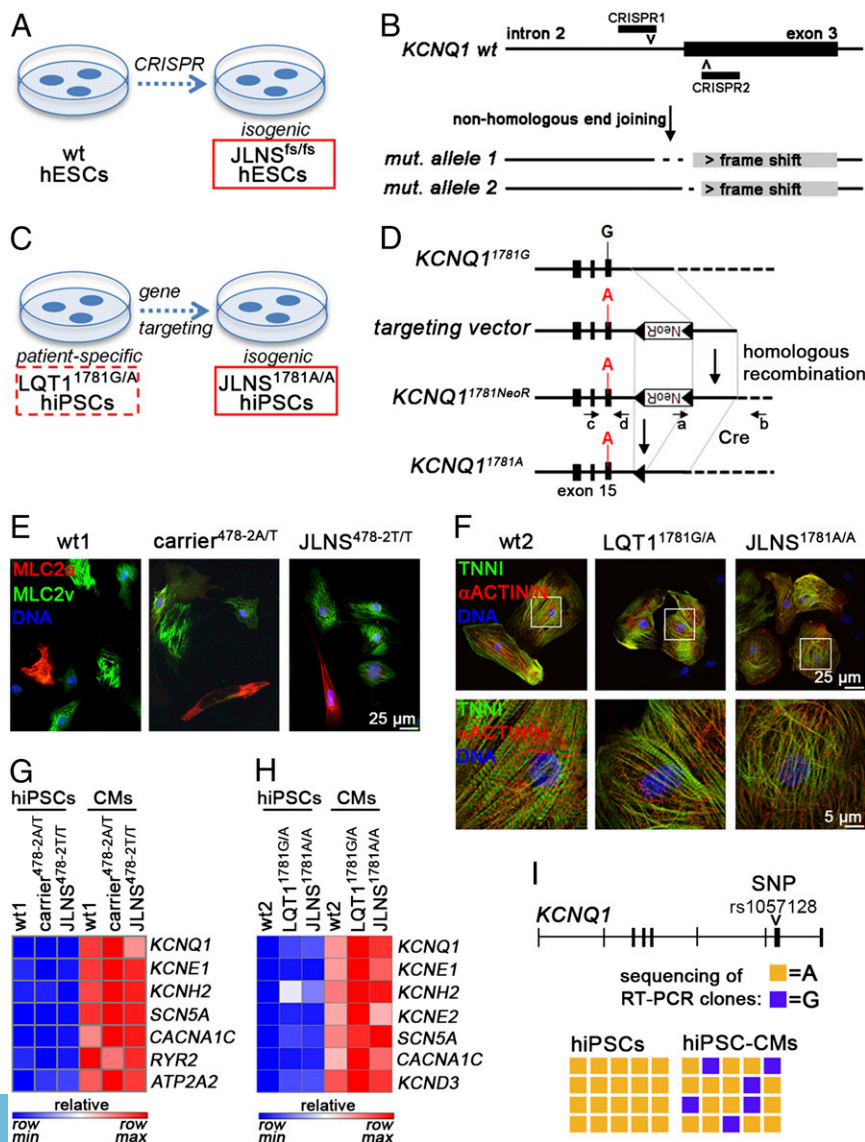


Fig. 2. Generation of isogenic JLNS hiPSC and hESC lines and general characterization of CMs. (A and B) CRISPR/Cas9-mediated homozygous disruption of the c.478-2 site in wt hESCs. Clones harboring frame shift-causing mutations on both alleles were isolated and expanded. (C and D) Schematic of the gene targeting strategy to introduce the c.1781G>A mutation in LQT1^{1781G/A} hiPSCs. Black boxes indicate exons (D). The wt allele of LQT1^{1781G/A} hiPSCs is shown with the black G in exon 15. The targeting vector has the mutation (A nucleotide, red) and a *loxP*-flanked G418-resistance cassette (*NeoR*). PCR primers a+b and c+d served to identify correctly targeted clones. (E and F) Immunofluorescence images of cardiac sarcomeric proteins MLC2a, MLC2v, TNNI, and α -actinin in the indicated lines following ~4 wk of in vitro differentiation and maturation. F, Lower is a magnification of the framed area in *Upper*. (G and H) Heat map representation of qPCR gene expression of cardiac ion channels, before and after cardiomyocyte differentiation. Data are relative to wt hiPSCs. (I) Loss of *KCNQ1* imprinting in wt1-CMs compared with undifferentiated hiPSCs. The SNP rs1057128 was used to discriminate between the two *KCNQ1* alleles in wt1 cells. Data denote sequencing results of 20 RT-PCR clones.

Electrophysiological Phenotype of JLNS^{478-2T/T} and carrier^{478-2A/T} hiPSC-CMs. We first characterized the electrophysiological properties of homozygous and heterozygous c.478-2A>T *KCNQ1* mutated hiPSC-CMs. Using multielectrode arrays (MEAs) that allow field potential duration (FPD) measurements in cell aggregates, JLNS^{478-2T/T}-CMs showed pronounced FPD prolongation compared with wt1 and carrier^{478-2A/T} CMs, although wt1- and carrier^{478-2A/T}-CMs did not differ significantly from each other (Fig. 3A). The same “recessive” manifestation of the electrophysiological phenotype was also observed in an independent set of (retrovirally) reprogrammed cell lines, ruling out cell clone-specific effects (Fig. S3A). To investigate the phenotype at the single-cell level, action potentials (APs) were measured by using the amphotericin-perforated patch-clamp technique. Fig. 3B and C show typical APs and average AP properties, respectively, measured at 1 Hz. AP duration at 20, 50, and 90% repolarization (APD₂₀, APD₅₀, and APD₉₀, respectively) in JLNS^{478-2T/T}-CMs were significantly increased compared with wt1 and heterozygous controls. In addition, AP amplitude (APA) and plateau amplitude (PlaA) were significantly increased in JLNS^{478-2T/T}-CMs. Maximal upstroke velocity (dV/dt_{max}) and resting membrane potential (RMP) were unchanged between cell lines. Of note, AP properties did not differ significantly between wt1- and carrier^{478-2A/T}-CMs. The

changes in AP duration between JLNS^{478-2T/T}-CMs and wt1- and carrier^{478-2A/T}-CMs were also detected at slower (0.5 Hz) and faster (2 Hz) pacing rates (Fig. 3D, Left). Interestingly, most of the JLNS-CMs could not be paced at 3 and 4 Hz (Fig. 3D, Right). These data are in agreement with the severe and recessive disease phenotype observed in JLNS.

Electrophysiological Phenotype of JLNS^{1781A/A} and LQT1^{1781G/A} hiPSC-CMs. Next, we investigated the electrophysiological properties of the c.1781G>A *KCNQ1* mutated CMs. Similarly to JLNS^{478-2T/T} CMs, FPD in homozygous JLNS^{1781A/A}-CMs was significantly increased compared with wt2-CMs. However, in contrast to the first set of hiPSC lines, the FPD in LQT1^{1781G/A}-CMs displayed an intermediate phenotype compared with wt2- and JLNS^{1781A/A}-CMs (Fig. 3E). APs measured at 1 Hz revealed significant AP prolongation in the JLNS^{1781A/A}-CMs compared with both LQT1^{1781G/A}- and wt2-CMs (Fig. 3F and G). Of note, and in agreement with MEA measurements, heterozygous LQT1^{1781G/A}-CMs displayed prolonged AP duration with respect to wt2-CMs. APA and PlaA were slightly increased in the JLNS^{1781A/A}- and LQT1^{1781G/A}-CMs compared with wt2-CMs, whereas dV/dt_{max} and RMP did not differ significantly between the lines. Significant differences in AP duration were also observed between the different hiPSC-CMs when they were stimulated at other

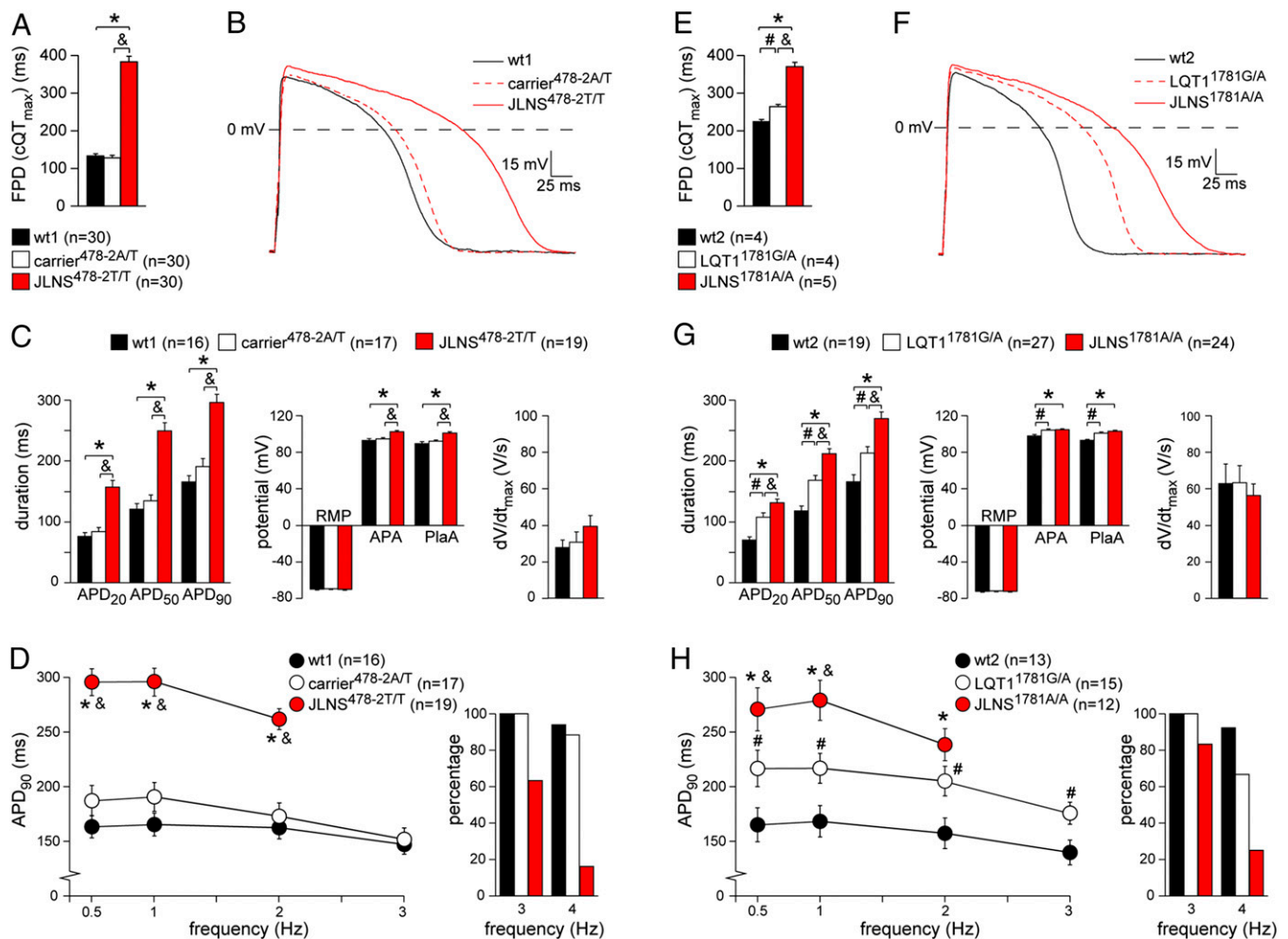


Fig. 3. Electrophysiological characterization of hiPSC-CMs. A–D show results for the wt1-, carrier^{478-2A/T}-, and JLNS^{478-2T/T}-CMs, and E–H for the wt2-, LQT1^{1781G/A}-, and JLNS^{1781A/A}-CMs. (A and E) FPD quantification on MEAs. (B and F) Representative APs at 1 Hz. (C and G) Average APD₂₀, APD₅₀, APD₉₀, dV/dt_{max}, RMP, APA, and PlaA. (D and H) Frequency dependence of APD₉₀ (Left). Most JLNS-CMs could not be paced at high frequencies (Right). Symbols *, &, and # indicate statistical significance based on pairwise comparisons (*P* < 0.05).

frequencies (0.5 and 2 Hz) (Fig. 3H, Left). Much like JLNS^{478-2T/T} CMs, fewer JLNS^{1781A/A}-CMs could be paced at 3 and 4 Hz compared with control CMs (Fig. 3H, Right). These results suggest a gene-dosage effect for the *KCNQ1* c.1781G>A mutation, in contrast to the strictly recessive c.478-2A>T nucleotide exchange. The more pronounced AP prolongation in the homozygous—compared with the heterozygous—state is in agreement with a more severe repolarization abnormality in JLNS than in LQT1.

The c.478-2A>T *KCNQ1* Mutation Abolishes I_{Ks} in Homozygous CMs. We next investigated the role of I_{Ks} in AP and FPD prolongation. Both *KCNQ1* and *KCNE1* were expressed at similar levels in the three cell lines (Fig. S3B). However, FPD in JLNS^{478-2T/T}-CMs was completely insensitive to the I_{Ks} blocker chromanol 293B (C293B) (Fig. 4A), whereas wt1- and carrier^{478-2A/T}-CMs showed FPD prolongation when I_{Ks} was blocked. Similarly, at the single-cell level, I_{Ks} was clearly detected in wt1- and carrier^{478-2A/T}-CMs, but virtually absent in JLNS^{478-2T/T}-CMs, as revealed by patch clamp electrophysiology using the I_{Ks} blocker JNJ303 (Fig. 4B). Furthermore, when E4031 was added to inhibit the rapid component of the delayed rectifier K^+ current, I_{Kr} , which cooperates with I_{Ks} in the repolarization of the AP, FPD was prolonged in

wt1- and carrier^{478-2A/T}-CMs without altering spontaneous beating, as expected (27) (Fig. S44). However, in JLNS^{478-2T/T}-CMs, E4031 reproducibly induced arrhythmia already at moderate doses (100 nM), in agreement with reduced repolarization reserve in these cells due to defective I_{Ks} . Collectively, these data indicate that the homozygous c.478-2A>T *KCNQ1* mutation is amorphic (nonfunctional).

The c.478-2A>T mutation Abolishes *KCNQ1* Protein Expression Through a Splicing Defect. We next sought to understand the molecular mechanisms underlying the recessive c.478-2A>T mutation. RT-PCR analysis showed that *KCNQ1* mRNA was smaller in JLNS^{478-2T/T}-CMs compared with wt1-CMs (Fig. 4C). Sequencing of the PCR product revealed skipping of exon 3 at the RNA level (Δ ex3), in agreement with the disruption of the intron 2 splice acceptor site at the DNA level. As expected, heterozygous carrier^{478-2A/T}-CMs expressed both alleles, which also confirmed the loss of *KCNQ1* imprinting (Fig. 4C and Fig. S4B). Interestingly, isoform-specific quantitative real-time PCR (qPCR) revealed that in carrier^{478-2A/T}-CMs, Δ ex3 *KCNQ1* was strongly diminished with respect to the wt transcript in the same cells, whereas the wt *KCNQ1* was expressed at levels comparable to wt1-CMs

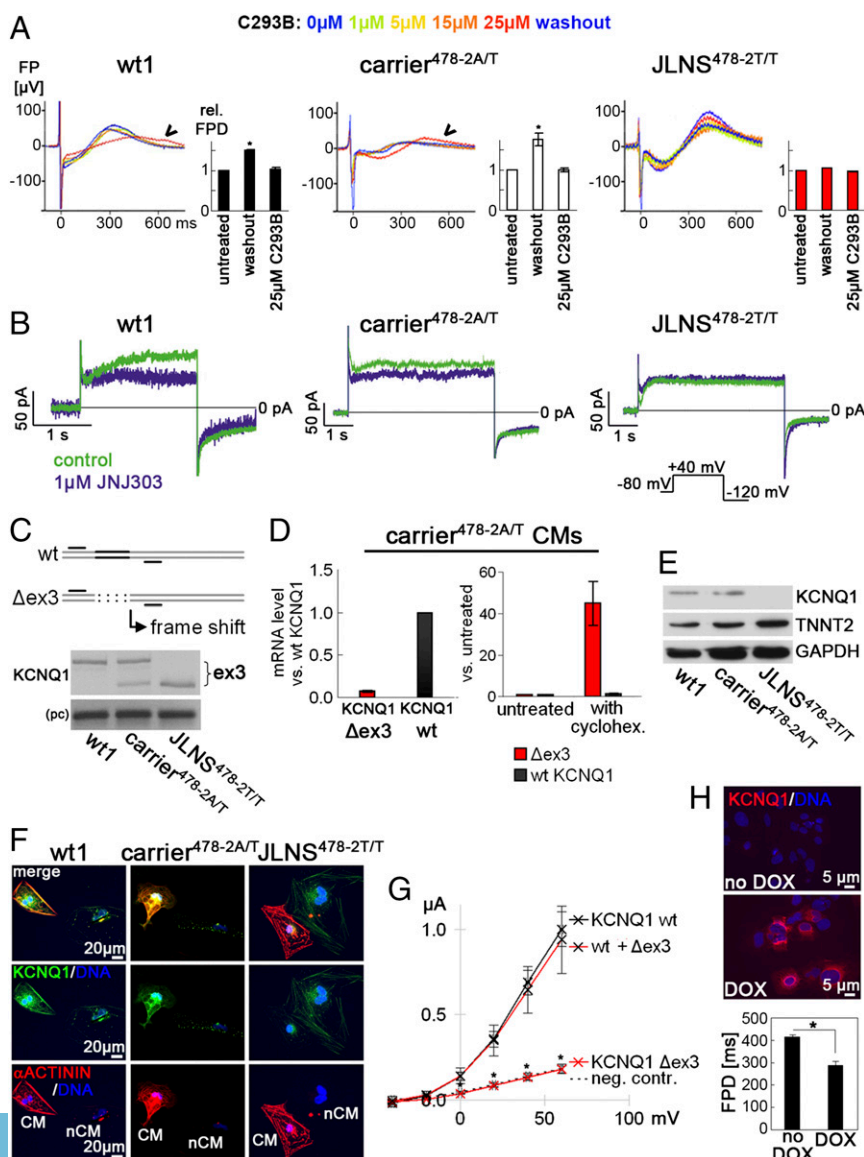


Fig. 4. Molecular mechanism underlying the c.478-2A/T mutation. (A) Effect of the I_{Ks} -blocker C293B on FPD. Single representative FPDs and average quantification are shown ($n = 3$). JLNS^{478-2T/T}-CMs are insensitive to the drug. (B) Representative current traces before (control) and after JNJ303 treatment in the indicated hiPSC-CMs. I_{Ks} is virtually absent in JLNS^{478-2T/T}-CMs. (Inset) Voltage protocol. (C) *KCNQ1* RT-PCR analysis in the indicated hiPSC-CMs. (Top) Schematic showing primers used to reveal the skipping of exon 3 in mutant hiPSC-CMs. (Bottom) Upper band, wt transcript; lower band, Δ ex3 mRNA; pc, positive control. (D) qPCR analysis of relative abundance of Δ ex3 and wt *KCNQ1* transcripts in carrier^{478-2A/T}-CMs under baseline conditions (Left) and upon cycloheximide treatment (Right); cycloheximide (30 μ g/mL, 3 h) selectively induces the Δ ex3 transcript. (E) Western blot analysis in hiPSC-CMs by using an antibody detecting the C terminus of *KCNQ1*. TNNT2 and GAPDH are shown as cardiac and loading controls, respectively. (F) Immunofluorescence analysis of *KCNQ1* and α -actinin in hiPSC-CMs. Noncardiomyocyte cells are shown next to CMs to indicate specificity of staining. (G) Average I-V relationship for I_{Ks} measured upon injection of wt, Δ ex3, and wt+ Δ ex3 mRNA into *Xenopus* oocytes ($n = 10$ each). (H) Genetic rescue of JLNS^{478-2T/T}-CMs by using DOX-inducible *KCNQ1* overexpression (Top); FPD shortening after 48 h of DOX treatment (Bottom, $n = 4$).

(Fig. 4D, Left and Fig. S4C). The exon 3 deletion causes a frame shift and premature stop codon in the *KCNQ1* ORF. The low Δex3 expression level may therefore be a result of nonsense-mediated mRNA decay (NMD) (28). To test this hypothesis, carrier^{478-2A/T}-CMs were transiently treated with cycloheximide, which inhibits the highly protein synthesis-dependent NMD pathway (29). Cycloheximide specifically up-regulated Δex3 - but not wt-*KCNQ1* mRNA in carrier^{478-2A/T}-CMs, suggesting that indeed, the mutant transcript is targeted by NMD (Fig. 4D, Right).

These data predict that the splice defect caused by the c.478-2A>T mutation abolishes *KCNQ1* protein in homozygous JLNS^{478-2T/T}-CMs, whereas in carrier^{478-2A/T} cells *KCNQ1* may be unaffected. Western blot and immunofluorescence analysis, using an antibody recognizing the C terminus of *KCNQ1*, confirmed this hypothesis (Fig. 4E and F). That the absence of *KCNQ1* correlates with significant FPD prolongation on MEAs was also evident in engineered isogenic JLNS^{fs/fs}-CMs, again indicating that the disease phenotype correlates with the loss of *KCNQ1* (Fig. S4D and E). Moreover, to test whether residual truncated *KCNQ1* could still exert a dominant negative effect on the wt channel, wt- and $\Delta\text{ex3-KCNQ1}$ mRNA isoforms were injected, alone and in combination, into *Xenopus* oocytes. Quantification of the corresponding currents confirmed complete loss of function of $\Delta\text{ex3-KCNQ1}$, without any dominant negative effect on wt-*KCNQ1* (Fig. 4G). Finally, to demonstrate functionally that AP and FPD prolongation in JLNS^{478-2T/T}-CMs was due to the absence of *KCNQ1*, a doxycycline (DOX)-inducible transgene was used to rescue the phenotype (Fig. S4F, Left). wt-*KCNQ1* transgene activation significantly reduced FPD prolongation and restored sensitivity to C293B (Fig. 4H and Fig. S4F, Right).

The c.1781G>A *KCNQ1* Mutation Reduces I_{Ks} in Heterozygous and Homozygous CMs. We then investigated the effect of the c.1781G>A *KCNQ1* mutation on I_{Ks} , measured as JNJ303-sensitive current (Fig. S5A). Fig. 5A and B show typical I_{Ks} and average JNJ303-sensitive tail current densities in wt2-, LQT1^{1781G/A}-, and JLNS^{1781A/A}-CMs. I_{Ks} tail current density was significantly smaller in JLNS^{1781A/A}-CMs compared with LQT1^{1781G/A}-CMs, whereas both mutated lines showed reduced I_{Ks} compared with wt2-CMs. By contrast, I_{Kr} tail density, measured as E4031-sensitive current, was comparable between the three lines (Fig. S5A and B), indicating a specific effect of the c.1781G>A mutation on I_{Ks} only. Taken together, these data suggest that the c.1781G>A mutation causes a larger reduction of I_{Ks} in the homozygous compared with the heterozygous genotype, in agreement with it being responsible for the more severe phenotype observed in JLNS^{1781A/A}-CMs (Fig. 3).

The c.1781G>A Mutation Results in a Partial *KCNQ1* Trafficking Defect. To investigate the molecular mechanisms underlying the electrophysiological phenotype of the c.1781G>A mutation, *KCNQ1* RNA and protein levels were determined in mutated and wt2-CMs. Channel expression at the protein and RNA levels was not reduced in either LQT1^{1781G/A}-CMs or in JLNS^{1781A/A}-CMs, ruling out a disease mechanism based on reduced *KCNQ1* abundance (Fig. 5C and Fig. S5C, Top). The β -subunit-encoding *KCNE1* was also expressed at similar levels in CMs of all lines, both at the protein and RNA levels (Fig. 5C, Top and Fig. S5C-E). To investigate the effect of the missense c.1781G>A mutation further, we examined ion channel expression by immunofluorescence. In all lines, *KCNQ1* was detected both on the cell surface and in intracellular compartments through which the channel normally transits during its biogenesis (Fig. 5D). However, both JLNS^{1781A/A}- and LQT1^{1781G/A}-CMs showed a reduction of *KCNQ1* at the outer cell membrane, with a more pronounced perinuclear signal in the majority of homozygous JLNS^{1781A/A}-CMs. These observations

suggest that the c.1781G>A mutation possibly interferes with the normal trafficking of the mutated channel to the cell membrane.

To confirm this result independently, we expressed GFP-tagged wt- or G1781A-mutated *KCNQ1* mRNA in *Xenopus* oocytes. Although the wt channel was abundant at the oocyte membrane, the mutated protein was greatly reduced in this compartment, although it was not completely absent (Fig. 5E). These data support a partial trafficking defect associated with the c.1781G>A *KCNQ1* mutation, as shown in other systems (30). To better understand the consequences of the R594Q amino acid exchange in the *KCNQ1* protein, we used a described in silico model (31). Molecular dynamic simulations suggested that the R594Q mutation may induce local structural rearrangements in the C-terminal domain of *KCNQ1*, with potential impact on tetrameric assembly (Fig. 5F and Fig. S5F). Taken together, these data suggest that the c.1781G>A mutation causes a partial *KCNQ1* trafficking defect, giving rise to LQT1 and JLNS phenotypes in a gene dosage-dependent manner.

JLNS-CMs Are Sensitive to β -Adrenergic and Proarrhythmic Stress but Benefit from β -Blockade and I_{Kr} Activation. Because JLNS patients are at high risk for arrhythmic events, we further examined the cellular models for effects of β -adrenergic stimulation and proarrhythmic drugs. APs were measured at 1 Hz in the absence or presence of 500 nM noradrenaline in CMs derived from all hiPSC lines. Typical APs and average effects on PlaA and APD₉₀ are shown in Fig. 6. β -adrenergic stimulation significantly increased the PlaA in all cases but only significantly prolonged the APD₉₀ in the JLNS^{478-2T/T}-, JLNS^{1781A/A}-, and LQT1^{1781G/A}-CMs. The increase in PlaA and in APD₉₀ was most pronounced in JLNS-CMs. These effects were partially reversed by treatment with the β -blocker propranolol (Fig. 6A and C). Neither early and delayed afterdepolarizations (EADs and DADs, respectively) nor spontaneous activity was observed at a pacing frequency of 1 Hz in the presence of noradrenaline.

To further assess whether mutated CMs were more sensitive to stress caused by proarrhythmic compounds not targeting *KCNQ1* itself, CMs from all lines were exposed to Cisapride, a gastrointestinal drug that has been withdrawn as a pharmaceutical because of arrhythmogenic side effects in humans (32, 33). Interestingly, although wt controls were rather insensitive to this drug, JLNS-CMs of both genotypes studied reproducibly developed arrhythmias at Cisapride doses of only 10–30 nM (Fig. 7A and B). Similar observations were made with engineered JLNS^{fs/fs}-CMs (Fig. S6A). Heterozygous CMs behaved like wt controls or showed an intermediate response, depending on the underlying mutation (Fig. 7A and B). Hence, these results revealed a strong correlation between genotype-dependent I_{Ks} reduction and risk of arrhythmic responses. Because I_{Kr} cooperates with I_{Ks} in the repolarization of cardiac AP, we investigated whether drug-based I_{Kr} activation could rescue disease phenotypes in mutated hiPSC-CMs. Indeed, NS1643, a small molecule that activates I_{Kr} by reducing hERG channel inactivation (34, 35), reduced the FPD in hiPSC-CMs (Fig. S6B) and effectively protected JLNS^{478-2T/T}-, JLNS^{1781A/A}-, and LQT1^{1781G/A}-CMs from Cisapride-induced arrhythmias (Fig. 7C and D and Fig. S6C).

Discussion

Reprogramming somatic cells from patients with genetic diseases has already advanced understanding of human maladies, particularly for diseases difficult to model in mice in vivo, or where primary tissues are difficult to access (36). Channelopathies have been particularly successfully modeled in hiPSC-CMs, in part because differing ion channel use in human and mouse heart often precludes direct extrapolation of phenotypes (37). Most striking has been their value in demonstrating the causality of specific LQTS mutations (20, 36). In the present study, we report a side-by-side analysis of two distinct *KCNQ1* mutations causing JLNS, the recessive and one of the most severe forms of LQTS

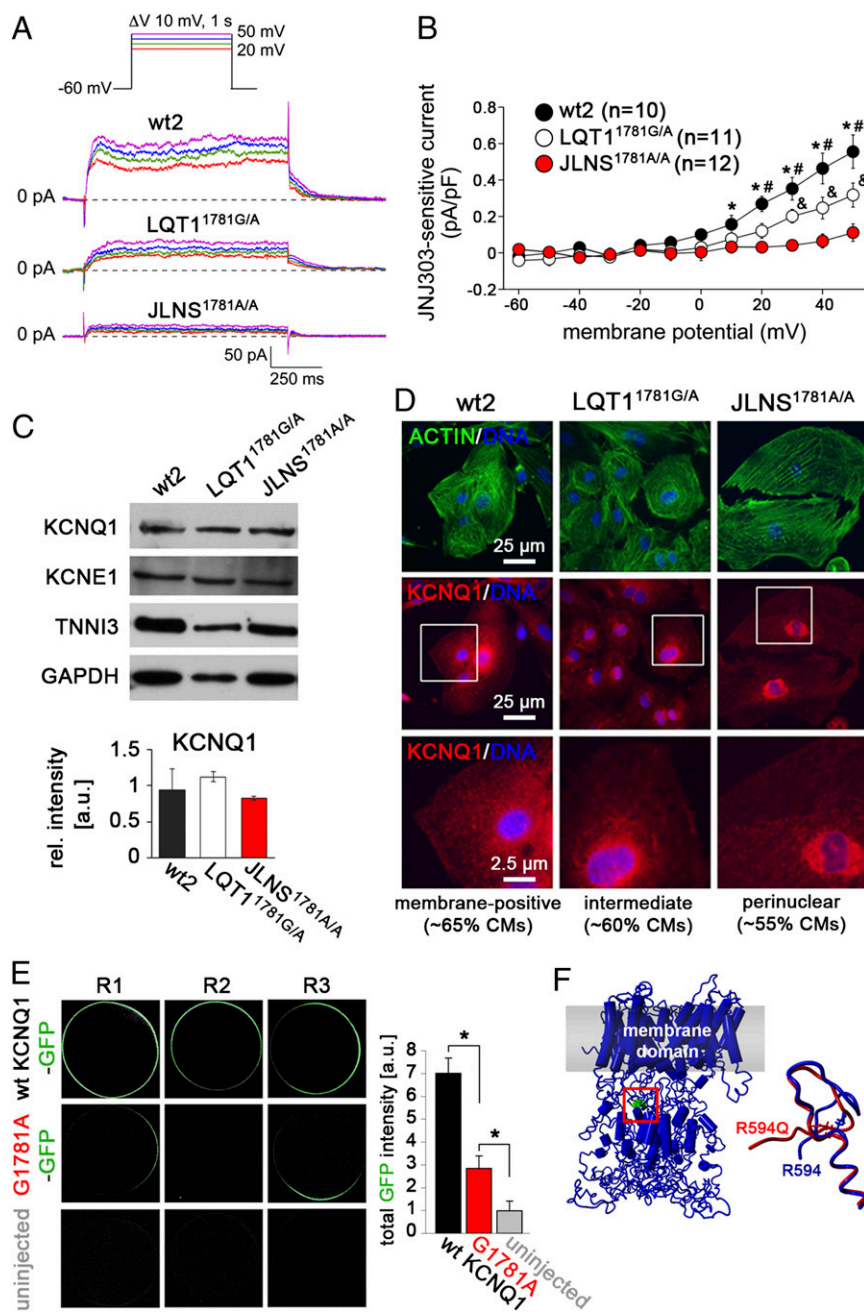


Fig. 5. Molecular mechanism underlying the c.1781G>A mutation. (A) Representative I_{Ks} measured as JNJ303-sensitive current in the indicated hiPSC-CMs. (Inset) Voltage protocol. (B) Average JNJ303-sensitive tail currents in hiPSC-CMs. Symbols *, &, and # indicate statistical significance based on pairwise comparisons ($P < 0.05$). (C) Western blot analysis of KCNQ1 and KCNE1 in hiPSC-CMs (Top). TNNI3 and GAPDH are used as cardiac and loading controls, respectively. (Bottom) Densitometric quantification of the KCNQ1 band, normalized to TNNI3 ($n = 2$). (D) Immunofluorescence analysis of KCNQ1 and actin in hiPSC-CMs. Bottom are a magnification of the area framed in Middle. Percentages of CMs showing the indicated staining patterns are indicated. (E) Representative confocal images showing membrane localization of KCNQ1 following injection of GFP-tagged wt and G1781A-mutated *KCNQ1* mRNA into *Xenopus* oocytes (Left). Image-based quantification of total GFP intensity at the outer cell membrane (Right, $n = 10$ each). (F) Structural model of KCNQ1 highlighting the position of the R594 residue in the C-terminal assembly domain (Left) and change in predicted local structure induced by the R594Q mutation (Right).

(2). We showed that both patient-derived and genetically engineered hiPSC-CMs recapitulated the aggressive cardiac phenotype of the disease, although the effect was mediated by different mechanisms.

Although previous JLNS studies in heterologous systems, where single ion channels are expressed in noncardiomyocytes, and mice have contributed significantly to understanding this disease, it has not always been possible to recapitulate the severity of cardiac phenotype caused by homozygous and heterozygous loss of function mutations in the same gene. Although the hearing loss observed in patients was recapitulated in *KCNQ1* or *KCNE1* knockout mice, clear cardiac phenotypes were not always evident (38, 39), illustrating the value of the human models here.

The c.478-2A>T mutation resulted in a splice defect at the mRNA level, causing exon 3 skipping and a reading frame shift (E160fs+138X). Consequently, KCNQ1 protein was not detectable

in JLNS^{478-2T/T}-CMs and, accordingly, these cells did not produce any I_{Ks} , providing an explanation for the severity of the observed cardiac phenotype. Indeed, the FPD was significantly shortened by transgenic *KCNQ1* overexpression in JLNS^{478-2T/T} cells. Moreover, in the heterozygous carrier^{478-2A/T}-CMs, Δ ex3 *KCNQ1* mRNA underwent NMD, whereas nonmutated *KCNQ1* mRNA was elevated above the expected 50% of the wt level. This latter observation may indicate a gene-dosage compensatory effect warranting further investigation. Furthermore, *Xenopus* oocyte assays revealed that Δ ex3 *KCNQ1* mRNA was nonfunctional and did not interfere with the wt form. In sum, these effects result in normal *KCNQ1* expression and function in heterozygous carrier^{478-2A/T}-CMs, explaining why the disease is recessive at the molecular and cellular level. This scenario likely has universal validity for nonsense and frame shift lesions in *KCNQ1* (10) and contrasts with dominant-negative effects of point mutations causing LQT1 (40, 41).

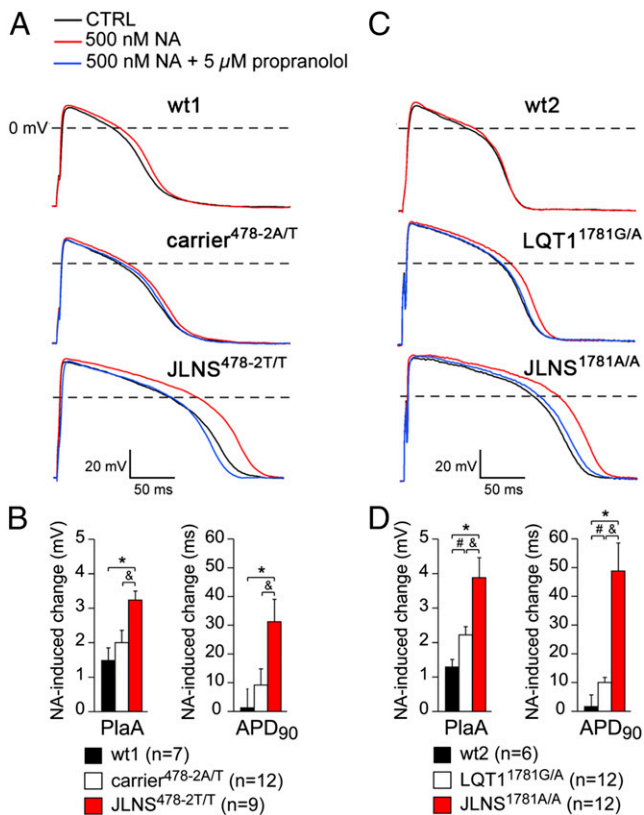


Fig. 6. Adrenergic stress and β -block in hiPSC-CMs. *A* and *B* show results for the wt1-, carrier^{478-2A/T}-, and JLNS^{478-2T/T}-CMs; and *C* and *D* for the wt2-, LQT1^{1781G/A}-, and JLNS^{1781A/A}-CMs. (*A* and *C*) Representative APs at 1 Hz in presence of noradrenaline (NA, red) or NA+propranolol (blue). (*B* and *D*) Average PlaA and APD₉₀ changes in presence of NA. Symbols *, &, and # indicate statistical significance based on pairwise comparisons ($P < 0.05$).

The c.1781G>A *KCNQ1* mutation, however, caused AP prolongation even when expressed heterozygously (LQT1^{1781G/A}-CMs), whereas the effect on AP prolongation was additive in the homozygous JLNS^{1781A/A}-CMs. This phenotype is in contrast with the strictly recessive loss-of-function phenotype caused by the c.478-2A>T *KCNQ1* lesion. These observations correspond well with clinical data from the patients from which the lines were derived: the heterozygous LQT1^{1781G/A} patient had a prolonged QT interval (QTc: 506 ms), whereas the carrier^{478-2A/T} had a normal ECG (QTc: 400 ms). Accordingly, molecular characterization of the c.1781G>A mutation revealed a distinct, gene dosage-dependent mechanism. Even as a heterozygote, the c.1781G>A point mutation reduced ion channel expression at the plasma membrane. This result could be a direct consequence of the localization of the c.1781G>A mutation in the C-terminal assembly domain of the protein (13) that might affect assembly of the mutated α -subunits and, consequently, the amount of protein reaching the membrane. Our results are in agreement with a recent study in which the same mutation was overexpressed in a heterologous system and a trafficking defect was evident (30, 42). In line with this mechanism, the homozygous mutation in JLNS^{1781A/A}-CMs caused a large reduction in I_{Ks} , albeit not a complete loss. The fact that in the homozygous mutant genotype some residual I_{Ks} current was present (~20%) would also support the possibility that homozygous patients harboring the c.1781G>A mutation could present with autosomal recessive LQTS (i.e., without deafness) (14). However, a larger number of such patients, together with their clinical data, would help provide a clear classification of this mutation. In

heterozygous LQT1^{1781G/A}-CMs, I_{Ks} was reduced by ~40%, indicating haploinsufficiency as the mechanism of action of this mutation, in agreement with a prolonged QT interval in our LQT1 patient. Of note, the lack of a dominant-negative effect is in agreement with the general mechanism proposed for the JLNS-causing mutations (11, 13, 43, 44).

JLNS and other LQTS subtypes are complex diseases in which phenotypes may be influenced by epigenetics and gene modifiers (44, 45). In addition, the type of mutation and its localization, and variable penetrance, have an impact on phenotype severity and complexity (43). These characteristics might explain the variability sometimes observed between patients even harboring the same mutation, and why individuals with heterozygous *KCNQ1* mutations can be either carriers or symptomatic patients (44). In line with these findings, engineered JLNS^{fs/fs} CMs showed strong but somewhat less pronounced FPD prolongation and arrhythmia phenotypes compared with their isogenic controls than patient-derived hiPSC-CMs compared with nonisogenic wt or heterozygous carrier CMs. In comparison, our data on the c.1781G>A *KCNQ1* mutation, where isogenic heterozygous and homozygous mutated CMs were compared, confirmed that precise gene targeting in hiPSC-CMs is a powerful tool to study the role of such mutations and evaluate dosage effects on disease severity (46).

One important characteristic of LQT1 and JLNS patients is their increased sensitivity to β -adrenergic triggers that are often the cause of sudden death (2, 47). Consistently, our data revealed significant AP prolongation in LQT1- and JLNS-CMs upon β -adrenergic stimulation, whereas AP prolongation was not observed in wt-CMs. These results are likely due to β -adrenergic-induced increases in L-type Ca^{2+} current (44) being counteracted by a prominent increase in I_{Ks} . The noradrenaline-induced AP prolongation in LQT1- and JLNS-CMs shows similarities with findings in freshly isolated ventricular CMs of human failing hearts (44), which have a reduced I_{Ks} (48).

Our data also revealed a particularly high sensitivity of JLNS-CMs to proarrhythmic stress. The increased susceptibility to drug-induced arrhythmias of JLNS^{1781A/A}- and JLNS^{478-2T/T}-CMs is likely a direct consequence of impaired I_{Ks} and concomitant reduction of the repolarization reserve (49). One promising application of hiPSC-CMs is screening for proarrhythmic side effects of certain drugs (50, 51). JLNS hiPSC-CMs may be particularly suited to this purpose because of their extraordinary sensitivity. It will be interesting to evaluate whether this effect holds true only for inhibitors of I_{Kr} —such as E4031 and Cisapride used here (32)—or if it also applies to other classes of cardiotoxic compounds.

Collectively, our data exemplify how distinct mechanisms may lead to similar JLNS phenotypes. These findings may also have implications for potential therapeutic strategies. For example, in the case of point mutations, the *KCNQ1* channel itself could in principle be considered as a drug target (52, 53), whereas in the case of frameshift mutations, it cannot. Moreover, LQT1 patients are often responsive to β -blocker therapy, but for the majority of JLNS patients, this treatment is only partially effective (2). Alternatively, compromised *KCNQ1* function might be compensated by drug-induced I_{Kr} activation (54). Using the hERG activator NS1643, FPDs were shortened in hiPSC-CMs, although this effect required high drug concentrations (30 μ M). Interestingly, the high susceptibility of JLNS-CMs to drug-induced arrhythmia could also be balanced by I_{Kr} activator treatment. This finding may also suggest alternative therapeutic perspectives for patients, although more potent compounds will be needed to make this strategy a realistic option.

Materials and Methods

Detailed procedures are outlined in *SI Materials and Methods*.

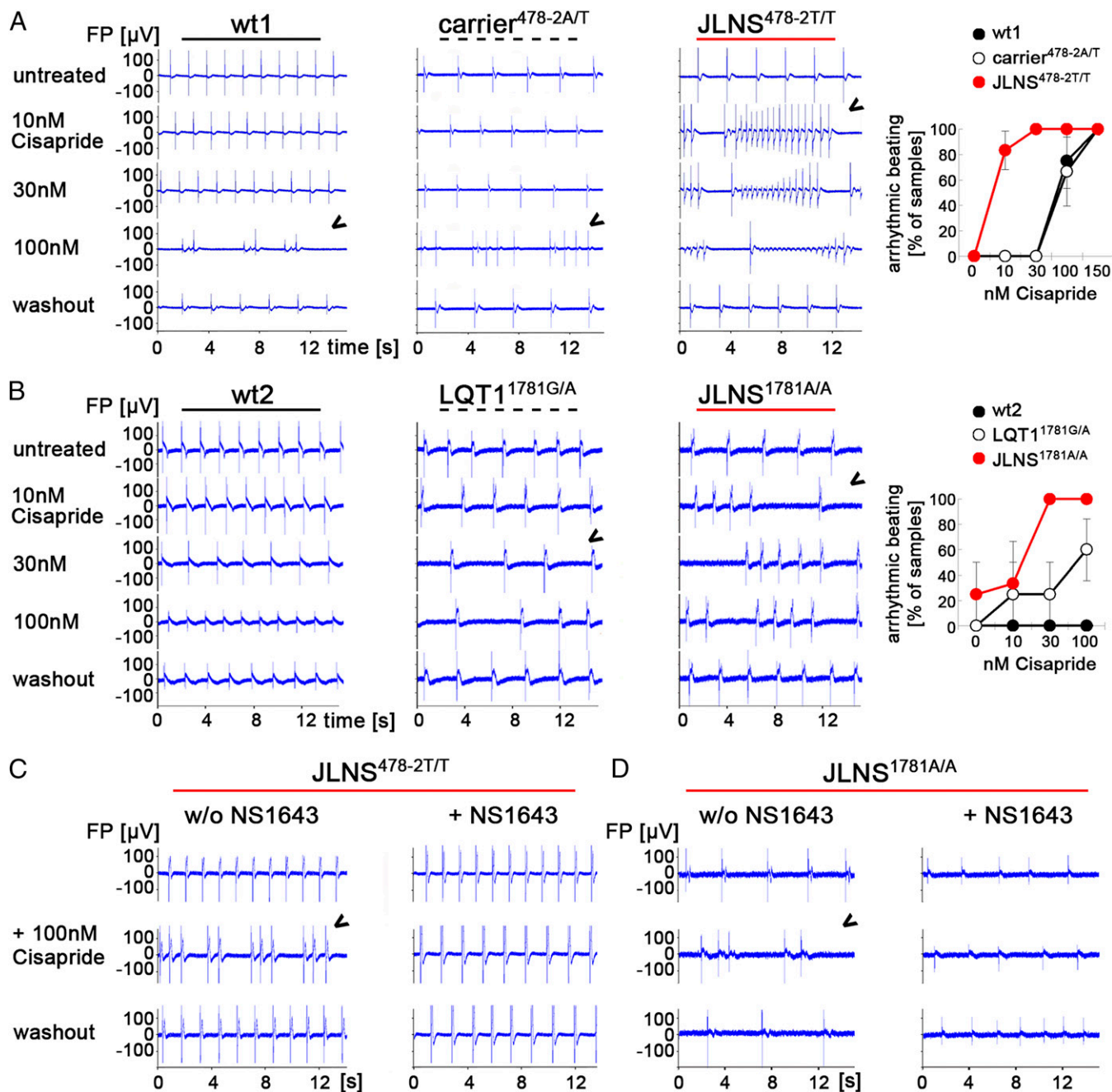


Fig. 7. Drug-induced arrhythmia phenotypes in JLNS-CMs and rescue by a hERG activator. (A and B) Cisapride causes arrhythmias in spontaneously beating wt⁻, carrier^{478-2A/T}, and JLNS^{478-2T/T}-CMs (A) as well as in LQT1^{1781G/A}- and JLNS^{1781A/A}-CMs. (B, Left) Representative MEA traces. Arrowheads indicate first appearance of arrhythmia. (B, Right) Statistics of arrhythmic beatings at different Cisapride doses from independent experiments ($n = 2-6$). (C and D) Pretreatment with the hERG channel activator NS1643 (30 μ M) protects JLNS^{478-2T/T}- (C) and JLNS^{1781A/A}-CMs (D) from high-dose Cisapride-induced arrhythmia. Representative MEA traces from independent experiments are shown.

Patients. The patients described in this study include the following: (i) a child with congenital deafness and QT interval prolongation (JLNS^{478-2T/T}, QTc: 570 ms; Fig. S1A), harboring the homozygous c.478-2A>T *KCNQ1* mutation—he is being treated with β -blockers and implantable cardioverter defibrillator implantation; (ii) his heterozygous and asymptomatic father (carrier^{478-2A/T}, QTc: 400 ms); and (iii) an independent woman presenting with LQT1 (LQT1^{1781G/A}, QTc: 506 ms; Fig. S1B) carrying the heterozygous c.1781G>A mutation in *KCNQ1*—she is asymptomatic but treated with β -blockers.

Cell Culture and Molecular Biology. hiPSC were generated, cultured, and differentiated into cardiomyocytes by using procedures specified in *SI Materials*

and Methods. Genetic engineering was achieved by using either a CRISPR/Cas-9 or a conventional homologous recombination strategy. Functional complementation was performed by using PiggyBac transposition. Sequencing, gene expression, and imprinting analysis were performed by using standard procedures. All primers used in this study are listed in Table S1. Immunofluorescence and Western blot analysis were performed according to standard procedures, using antibodies given in *SI Materials and Methods*.

Cellular Electrophysiology. Experimental details about FPD, APs, I_{Ks} , and I_{Kr} measurements and analysis in hiPSC-CMs and *Xenopus laevis* oocytes are described in *SI Materials and Methods*.

Statistics. Data are presented as mean \pm SEM. Groups were compared by using one-way ANOVA followed by Tukey's test or two-way repeated measures ANOVA followed by pairwise comparison by using the Student–Newman–Keuls test. MEAs data were analyzed on the basis of two-sided unpaired Student *t* tests. *P* < 0.05 was considered statistically significant and is indicated by *, &, and #.

ACKNOWLEDGMENTS. We thank M. Ohtaka, K. Nishimura, and M. Nakanishi (National Institute of Advanced Industrial Science and Technology, Japan) for providing the Sendai virus and reprogramming protocol; the hiPSC core facility (Leiden University Medical Center) for hiPSCs spontaneous differentiation; F. Stewart (Technical University of Dresden) for providing the R6K plasmid; M. J. M. van der Burg, K. Suzhai, H. Tanke (Leiden University Medical

Center) for karyotyping; M. J. Goumans (Leiden University Medical Center) for providing adult heart samples; K. Adachi (Max Planck Institute Münster) for providing the KA0717 vector; and A. Wilde and N. Hofman (both Academic Medical Center Amsterdam) for constructive discussion. We are especially thankful to all the patients for tissue donation. This work was supported by European Research Council Grant 323182 (to C.L.M.); ZonMW Animal Alternatives Grant 114000101 (to C.L.M.); Netherlands Proteomics Consortium Grant 050-040-250 (to C.L.M.); the Netherlands Institute of Regenerative Medicine (C.L.M. and C.D.); European Union Marie Curie FP7-people-2011-IEF Programme, HPSCLQT 29999 (to M.B.) and a short-term EMBO Fellowship, ASTF 387.00-2011 (to C.D.); Foundation Leducq (E.S.-B.), Interdisciplinary Center for Clinical Research (E.S.-B.); Eva Luise and Horst Köhler Foundation (B.G.); Bundesinstitut für Risikobewertung, FK-3-1329-471 (to B.G.); and Chemical Genomics Centre of the Max Planck Society (B.G.).

- Jervell A, Lange-Nielsen F (1957) Congenital deaf-mutism, functional heart disease with prolongation of the Q-T interval and sudden death. *Am Heart J* 54(1):59–68.
- Schwartz PJ, et al. (2006) The Jervell and Lange-Nielsen syndrome: Natural history, molecular basis, and clinical outcome. *Circulation* 113(6):783–790.
- Jespersen T, Grunnet M, Olesen SP (2005) The KCNQ1 potassium channel: From gene to physiological function. *Physiology (Bethesda)* 20:408–416.
- Schulze-Bahr E, et al. (1997) KCNE1 mutations cause Jervell and Lange-Nielsen syndrome. *Nat Genet* 17(3):267–268.
- Romano C, Gemme G, Pongiglione R (1963) [Rare cardiac arrhythmias of the pediatric age. II. syncopal attacks due to paroxysmal ventricular fibrillation. (Presentation of 1st case in Italian pediatric literature)]. *Clin Pediatr (Bologna)* 45:656–683.
- Ward OC (1964) A New Familial Cardiac Syndrome in Children. *J Ir Med Assoc* 54:103–106.
- Schwartz PJ, Ackerman MJ, George AL, Jr, Wilde AA (2013) Impact of genetics on the clinical management of channelopathies. *J Am Coll Cardiol* 62(3):169–180.
- Splawski I, et al. (2004) Ca(V)_{1.2} calcium channel dysfunction causes a multisystem disorder including arrhythmia and autism. *Cell* 119(1):19–31.
- Crotti L, et al. (2013) Calmodulin mutations associated with recurrent cardiac arrest in infants. *Circulation* 127(9):1009–1017.
- Splawski I, et al. (2000) Spectrum of mutations in long-QT syndrome genes. KVLQT1, HERG, SCN5A, KCNE1, and KCNE2. *Circulation* 102(10):1178–1185.
- Tyson J, et al. (2000) Mutational spectrum in the cardioauditory syndrome of Jervell and Lange-Nielsen. *Hum Genet* 107(5):499–503.
- Jackson H, et al. (2014) LQTS in Northern BC: Homozygosity for KCNQ1 V205M presents with a more severe cardiac phenotype but with minimal impact on auditory function. *Clin Genet* 86(1):85–90.
- Schmitt N, et al. (2000) A recessive C-terminal Jervell and Lange-Nielsen mutation of the KCNQ1 channel impairs subunit assembly. *EMBO J* 19(3):332–340.
- Bhuiyan ZA, Wilde AA (2013) IKs in heart and hearing, the ear can do with less than the heart. *Circ Cardiovasc Genet* 6(2):141–143.
- Neyroud N, et al. (1997) A novel mutation in the potassium channel gene KVLQT1 causes the Jervell and Lange-Nielsen cardioauditory syndrome. *Nat Genet* 15(2):186–189.
- Splawski I, Timothy KW, Vincent GM, Atkinson DL, Keating MT (1997) Molecular basis of the long-QT syndrome associated with deafness. *N Engl J Med* 336(22):1562–1567.
- Tranebjærg L, Bathen J, Tyson J, Bitner-Glindzic M (1999) Jervell and Lange-Nielsen syndrome: A Norwegian perspective. *Am J Med Genet* 89(3):137–146.
- Inoue H, Nagata N, Kurokawa H, Yamanaka S (2014) iPS cells: A game changer for future medicine. *EMBO J* 33(5):409–417.
- Moretti A, Laugwitz KL, Dorn T, Sinnecker D, Mummery C (2013) Pluripotent stem cell models of human heart disease. *Cold Spring Harb Perspect Med* 3(11).
- Müller M, Seufferlein T, Illing A, Homann J (2013) Modelling human channelopathies using induced pluripotent stem cells: A comprehensive review. *Stem Cells Int* 2013:496501.
- Ma D, et al. (2013) Modeling type 3 long QT syndrome with cardiomyocytes derived from patient-specific induced pluripotent stem cells. *Int J Cardiol* 168(6):5277–5286.
- Huang L, Bitner-Glindzic M, Tranebjærg L, Tinker A (2001) A spectrum of functional effects for disease causing mutations in the Jervell and Lange-Nielsen syndrome. *Cardiovasc Res* 51(4):670–680.
- Nishimura K, et al. (2011) Development of defective and persistent Sendai virus vector: A unique gene delivery/expression system ideal for cell reprogramming. *J Biol Chem* 286(6):4760–4771.
- Müller FJ, et al. (2011) A bioinformatic assay for pluripotency in human cells. *Nat Methods* 8(4):315–317.
- Ran FA, et al. (2013) Double nicking by RNA-guided CRISPR Cas9 for enhanced genome editing specificity. *Cell* 154(6):1380–1389.
- Lee MP, Hu RJ, Johnson LA, Feinberg AP (1997) Human KVLQT1 gene shows tissue-specific imprinting and encompasses Beckwith-Wiedemann syndrome chromosomal rearrangements. *Nat Genet* 15(2):181–185.
- Braam SR, et al. (2010) Prediction of drug-induced cardiotoxicity using human embryonic stem cell-derived cardiomyocytes. *Stem Cell Res (Amst)* 4(2):107–116.
- Maquat LE (2004) Nonsense-mediated mRNA decay: Splicing, translation and mRNP dynamics. *Nat Rev Mol Cell Biol* 5(2):89–99.
- Carter MS, et al. (1995) A regulatory mechanism that detects premature nonsense codons in T-cell receptor transcripts in vivo is reversed by protein synthesis inhibitors in vitro. *J Biol Chem* 270(48):28995–29003.
- Harmer SC, et al. (2014) Cellular mechanisms underlying the increased disease severity seen for patients with long QT syndrome caused by compound mutations in KCNQ1. *Biochem J* 462(1):133–142.
- Strutz-Seebohm N, et al. (2011) Structural basis of slow activation gating in the cardiac I_{Ks} channel complex. *Cell Physiol Biochem* 27(5):443–452.
- Rampe D, Roy ML, Dennis A, Brown AM (1997) A mechanism for the proarrhythmic effects of cisapride (Propulsid): High affinity blockade of the human cardiac potassium channel HERG. *FEBS Lett* 417(1):28–32.
- Bran S, Murray WA, Hirsch IB, Palmer JP (1995) Long QT syndrome during high-dose cisapride. *Arch Intern Med* 155(7):765–768.
- Hansen RS, et al. (2006) Activation of human ether-a-go-go-related gene potassium channels by the diphenylurea 1,3-bis-(2-hydroxy-5-trifluoromethyl-phenyl)-urea (NS1643). *Mol Pharmacol* 69(1):266–277.
- Casis O, Olesen SP, Sanguinetti MC (2006) Mechanism of action of a novel human ether-a-go-go-related gene channel activator. *Mol Pharmacol* 69(2):658–665.
- Bellini M, Marchetto MC, Gage FH, Mummery CL (2012) Induced pluripotent stem cells: The new patient? *Nat Rev Mol Cell Biol* 13(11):713–726.
- Davis RP, et al. (2012) Cardiomyocytes derived from pluripotent stem cells recapitulate electrophysiological characteristics of an overlap syndrome of cardiac sodium channel disease. *Circulation* 125(25):3079–3091.
- Rivas A, Francis HW (2005) Inner ear abnormalities in a Kcnq1 (Kvlqt1) knockout mouse: A model of Jervell and Lange-Nielsen syndrome. *Otol Neurotol* 26(3):415–424.
- Casimiro MC, et al. (2001) Targeted disruption of the Kcnq1 gene produces a mouse model of Jervell and Lange-Nielsen Syndrome. *Proc Natl Acad Sci USA* 98(5):2526–2531.
- Egashira T, et al. (2012) Disease characterization using LQTS-specific induced pluripotent stem cells. *Cardiovasc Res* 95(4):419–429.
- Moretti A, et al. (2010) Patient-specific induced pluripotent stem-cell models for long-QT syndrome. *N Engl J Med* 363(15):1397–1409.
- Wilson AJ, Quinn KV, Graves FM, Bitner-Glindzic M, Tinker A (2005) Abnormal KCNQ1 trafficking influences disease pathogenesis in hereditary long QT syndromes (LQT1). *Cardiovasc Res* 67(3):476–486.
- Moss AJ, et al. (2007) Clinical aspects of type-1 long-QT syndrome by location, coding type, and biophysical function of mutations involving the KCNQ1 gene. *Circulation* 115(19):2481–2489.
- Saenen JB, Vrints CJ (2008) Molecular aspects of the congenital and acquired Long QT Syndrome: Clinical implications. *J Mol Cell Cardiol* 44(4):633–646.
- Seebohm G, et al. (2008) Long QT syndrome-associated mutations in KCNQ1 and KCNE1 subunits disrupt normal endosomal recycling of IKs channels. *Circ Res* 103(12):1451–1457.
- Bellini M, et al. (2013) Isogenic human pluripotent stem cell pairs reveal the role of a KCNH2 mutation in long-QT syndrome. *EMBO J* 32(24):3161–3175.
- Schwartz PJ, et al. (2001) Genotype-phenotype correlation in the long-QT syndrome: Gene-specific triggers for life-threatening arrhythmias. *Circulation* 103(1):89–95.
- Janse MJ (2004) Electrophysiological changes in heart failure and their relationship to arrhythmogenesis. *Cardiovasc Res* 61(2):208–217.
- Braam SR, et al. (2013) Repolarization reserve determines drug responses in human pluripotent stem cell derived cardiomyocytes. *Stem Cell Res (Amst)* 10(1):48–56.
- Rajamohan D, et al. (2013) Current status of drug screening and disease modelling in human pluripotent stem cells. *BioEssays* 35(3):281–298.
- Navarrete EG, et al. (2013) Screening drug-induced arrhythmia [corrected] using human induced pluripotent stem cell-derived cardiomyocytes and low-impedance microelectrode arrays. *Circulation* 128(11, Suppl 1):S3–S13.
- Xu X, et al. (2002) Increasing I(Ks) corrects abnormal repolarization in rabbit models of acquired LQTS and ventricular hypertrophy. *Am J Physiol Heart Circ Physiol* 283(2):H664–H670.
- Nattel S, Carlsson L (2006) Innovative approaches to anti-arrhythmic drug therapy. *Nat Rev Drug Discov* 5(12):1034–1049.
- Zhang H, et al. (2012) Modulation of hERG potassium channel gating normalizes action potential duration prolonged by dysfunctional KCNQ1 potassium channel. *Proc Natl Acad Sci USA* 109(29):11866–11871.