GENERAL ARTICLE

Genetic variants in TRPM7 associated with unexplained stillbirth modify ion channel function

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Abstract

Stillbirth is the loss of a fetus after 22 weeks of gestation, of which almost half go completely unexplained despite post-mortem. We recently sequenced 35 arrhythmia-associated genes from 70 unexplained stillbirth cases. Our hypothesis was that deleterious mutations in channelopathy genes may have a functional effect in utero that may be pro-arrhythmic in the developing fetus. We observed four heterozygous, nonsynonymous variants in transient receptor potential melastatin 7 (TRPM7), a ubiquitously expressed ion channel known to regulate cardiac development and repolarization in mice. We used site-directed mutagenesis and single-cell patch-clamp to analyze the functional effect of the four stillbirth mutants on TRPM7 ion channel function in heterologous cells. We also used cardiomyocytes derived from human pluripotent stem cells to model the contribution of TRPM7 to action potential morphology. Our results show that two TRPM7 variants, p.G179V and p.T860M, lead to a marked reduction in ion channel conductance. This observation was underpinned by a lack of measurable TRPM7 protein expression, which in the case of p.T860M was due to rapid proteasomal degradation. We also report that human hiPSC-derived cardiomyocytes possess measurable TRPM7 currents; however, siRNA knockdown did not directly affect action potential morphology. TRPM7 variants found in the unexplained stillbirth population adversely affect ion channel function and this may precipitate fatal arrhythmia in utero.

Introduction

Stillbirth is the devastating loss of a fetus after 22 weeks of gestation during pregnancy. In the UK, there were 4.64 stillbirths per 1000 total births in 2013 (1). The majority of these are caused by obstetric complications, placental insufficiency, genetic disorders, infection and umbilical cord anomalies. However, despite extensive post-mortem investigation, 15–45% of stillbirths go completely unexplained (1–3).

Deleterious mutations within ion channel genes can cause severe arrhythmias and are known causes of sudden death in young adults (4). The cause of death is usually found after a negative post-mortem. We recently sequenced a custom panel...
of 35 arrhythmia-associated genes in 70 unexplained stillbirth cases from the Cardiac Ion Channelopathies in Stillbirth study (5). We found four cases to have putative pathogenic variants in four long QT syndrome genes, KCNQ1, KCNQ2, KCNS1, and KCNJ2. We subsequently characterized the functional effect of the heterozygous variant (p.R40Q) found in the KCNJ2 gene. This gene encodes the inwardly rectifying potassium channel Kir2.1, which controls the resting membrane potential and contributes to cardiac repolarization (Phase 3) (6). The p.R40Q mutation led to significantly reduced inward currents compared to wild-type (WT) channels, which would be predicted to have a deleterious effect on the cardiomyocyte ability to regulate its membrane voltage. Alongside these data, we observed predicted damaging variants in several other genes, including transient receptor potential melastatin 7 (TRPM7), a gene implicated from genome-wide association studies of QT interval (7). Four heterozygous nonsynonymous variants were observed in TRPM7 in four individual cases.

TRPM7 is a non-selective cation channel with a serine/threonine kinase domain in the C-terminus and is ubiquitously expressed in mammalian cells (8). Current opinion regards TRPM7 as a key transporter of divalent cations (Zn\(^{2+}\), Mg\(^{2+}\) and Ca\(^{2+}\)) across the cell membrane (9,10), while the \(\alpha\)-kinase is capable of phosphorylating downstream effector kinases and modifying chromatin (11,12). TRPM7 plays a key role in dictating cell proliferation, survival, apoptosis alongside organogenesis and embryonic survival (13–16). A previously reported genetic variant, p.T1482I, has been shown to increase the sensitivity of TRPM7 to inhibition by Mg\(^{2+}\) (17). Temporal control of TRPM7 expression in the mouse embryo is required for correct cardiac development (15).

The four nonsynonymous variants in TRPM7 lie within conserved amino acid positions and were predicted to functionally affect the protein at least one of three predicted software tools (PolyPhen, SIFT or Mutation Taster (18,19)). The first two, p.G179V and p.R494Q, lie within the N-terminal domain of TRPM7. The third, p.T860M, is within the second transmembrane domain, whilst the final variant, p.E1205G, resides in the C-terminal domain, upstream of the alpha kinase (Table 1).

In this study, we transfected a TRPM7 expressing plasmid into HEK293 and CHO-K1 cells and record the resulting current using whole-cell patch-clamp electrophysiology. We demonstrate that three TRPM7 variants (p.G179V, p.R494Q and p.T860M) significantly alter current densities in comparison to the WT ion channel. This effect was not due to an altered response to magnesium inhibition nor was TRPM7 transcription perturbed. Instead, we found a lack of protein expression in p.G179V and p.T860M-transfected cells, alongside an increase in p.R494Q protein. Interestingly, inhibiting the 26S proteasome with MG132 treatment increased p.R494Q expression and inhibited p.T860M expression. To further characterize the importance of TRPM7 in developing cardiomyocytes, we measured TRPM7 current in hiPSC-derived cardiomyocytes. While we found TRPM7 expression to increase in the early stages of differentiation, and there is a large outward TRPM7-like current, knockdown with TRPM7 siRNA had no effect on cardiomyocyte action potential morphology.

### Results

### Expressing TRPM7 in HEK293 and CHO-K1 cells

To ascertain the effect the nonsynonymous mutations may have on TRPM7 function, we transfected both CHO-K1 and HEK293 cells with a WT TRPM7 expression plasmid. After 5–10 min of dialysis, large outward currents were detected, with an I–V relationship typical of TRPM7 (Fig. 1A). We analyzed outward and inward current at +80 and –80 mV, respectively, in both CHO-K1 and HEK293 cells (Fig. 1B and C). Using CHO-K1 cells, we demonstrated that these currents were susceptible to inhibition by Mg\(^{2+}\) (17). HEK293 cell TRPM7 current was also inhibited by treatment with extracellular magnesium (data not shown). These data confirmed that we had an efficient expression system to test the effect of our stillbirth sequenced variants on TRPM7 function.

### Variants alter TRPM7 current density in CHO-K1 and HEK293 cells

We transfected both cell lines with the four variants of TRPM7 and recorded the current–voltage relationships. Due to its likely endogenous expression in both cell lines, we found TRPM7 I–V relationships in all cells (Fig. 2). However, when analyzing grouped current density data, we found a significant reduction in current density in CHO-K1 cells when transfected with p.G179V (P < 0.05, Fig. 2A). We also saw a significant increase in current density at both +80 and –80 mV when expressing p.R494Q compared to WT–transfected cells (P < 0.001, Fig. 2A). When these experiments were repeated in HEK293 cells, we found that both p.G179V and p.T860M current density was markedly reduced compared to WT TRPM7 (P < 0.01) with a trend to reduction for E1205G (Fig. 2B). However, in comparison to CHO-K1 expression of p.R494Q, current density was equal between WT and p.R494Q expressing cells (P = 0.98, Fig. 2B).

### Table 1. Summary of TRPM7 variants found in stillbirth cases. We sequenced four predicted damaging variants represented as both cDNA (c.) and protein (p.) changes. Listed are their relative locations within known domains of the TRPM7 protein. GERP++ score estimates the level of functional constraint that DNA position is under, within coding regions the average conservation score is 2 (28). PolyPhen and SIFT scores rate amino acid substitutions on their probability of impacting protein structure and function by providing values between 0 and 1 (18,19). Variants rated above 0.85 by PolyPhen are confidently predicted to be damaging, while those rated below 0.05 by SIFT are considered likely to be deleterious. Mutation taster analyses substitutions based upon base conservation, specific amino acid substitutions, functional domain location and other possible splicing effects. All TRPM7 variants were rated as 1 and annotated as disease causing, gnomAD frequency represents the variant’s abundance in the online genome aggregation database project, containing 125,748 exome sequences.

<table>
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<th>Mutation</th>
<th>Domain</th>
<th>GERP++</th>
<th>PolyPhen</th>
<th>Mutation taster</th>
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Figure 1. TRPM7 current recorded from transfected CHO-K1 and HEK293 cells. (A) Representative TRPM7 current trace from CHO-K1 cell transfected with TRPM7 at initial break-in ($t = 0$ s) and after current run-up ($t = 500$ s). (B) Current density at $+80$ mV from both CHO-K1 and HEK293 cells transiently transfected with TRPM7 (red) compared with untransfected cells (black). (C) Matching inward current density from cells in (A), analyzed at $-80$ mV. (D) Whole-cell current trace before (black) and after (red) addition of 10 mM magnesium to extracellular solution during whole-cell recording of a WT TRPM7-transfected CHO-K1 cell. (E) Time course of CHO-K1 whole-cell current showing immediate inhibition following 10 mM magnesium treatment. (F) Summary of whole-cell peak current data and following 10 mM magnesium treatment in both WT TRPM7-transfected and UT CHO-K1 cells at $+80$ mV. (G) Whole-cell current trace before (black) and after (red) addition of 50 μM 2-APB to extracellular solution during whole-cell recording of a WT TRPM7-transfected CHO-K1 cell. (H) Time course of current at $+80$ mV from cell shown in (G), showing current reduction over 300 s following 2-APB addition. WT, UT = untransfected. *$P < 0.05$, **$P < 0.001$. 

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Reducing extracellular Mg$^{2+}$ amplifies difference between WT and mutant TRPM7 channels

We wanted to investigate whether the stillbirth mutations were affecting the TRPM7 channel’s response to magnesium inhibition. We perfused transfected CHO-K1 cells with extracellular solution supplemented with increasing concentrations of magnesium from 0.5 to 6 mM after allowing current recordings to plateau. In these experiments, we found a significant decrease in current density at +80 mV in cells transfected with either p.G179V or p.T860M TRPM7 compared to WT channels, but only at 0 mM Mg$^{2+}$ (P < 0.01 and 0.05, respectively, Fig. 3A and C). However, when bath magnesium was increased, any difference in Mg$^{2+}$-dependent inhibition between these variants and WT TRPM7 was attenuated and eventually abolished. Cells expressing p.R494Q TRPM7 showed a significant increase in current density at +80 mV compared to WT cells at both 0 and 0.5 mM Mg$^{2+}$ (P < 0.001, Fig. 3B). We calculated the IC$_{50}$ for WT-transfected cells at 0.62 mM Mg$^{2+}$ with a Hill coefficient of −1.21. Further analysis of stillbirth variant-transfected cell IC$_{50}$ and Hill coefficient values was not significantly different to WT TRPM7 (Supplementary Material, Table S3). Interestingly, the inward current for this variant analyzed at −80 mV was increased compared to WT TRPM7 at 0, 0.5 and 1 mM Mg$^{2+}$ (P < 0.001, 0.001 and 0.05, respectively, Fig. 3E). Suggesting that despite physiological levels of magnesium, this variant increased...
both inward and outward ionic conductance compared to the WT TRPM7 channel. These data also suggest that any difference in whole-cell TRPM7 ion channel conductance are not the result of a variant influencing the channel's response to magnesium inhibition but are more likely due to alterations in expression.

In summary, our electrophysiological data show that G179V and T860M result in a loss of channel function and R494Q, a gain of function, which was particularly apparent in conditions of low extracellular Mg2+.

Effect of variants on TRPM7 protein and mRNA expression in heterologous cells

To investigate whether this decrease in current density was due to altered protein levels, whole cell lysates were harvested from transfected cells and western blotting was performed. G179V and T860M TRPM7 mutants did not express well in transiently transfected cells (Fig. 4A). Densitometry measurements showed a significant decrease in HEK293 cells transfected with either p.G179V or p.T860M TRPM7 compared to WT TRPM7. However, when we investigated TRPM7 and mutant mRNA levels in these cells using qPCR, the mRNA expression level was similar (Fig. 4C).

To test whether this reduction in protein was due to rapid degradation, we harvested cell lysates after overnight treatment with the proteasomal inhibitor MG132. We observed an increase in p.R494Q-transfected cells compared to WT, and we may have able to detect low level expression of p.T860M TRPM7 protein (Supplementary Material, Fig. S1).

Confirming the presence of TRPM7 current in hiPSC-derived cardiomyocytes

We used an in vitro model of cardiomyocyte generation using hiPSC cells to first ascertain TRPM7 expression in human cardiac cells and to test the effect that reduced TRPM7 expression may have on the electrophysiological properties of cardiomyocytes. We used an in-house hiPSC line (named HSIM) and differentiated the cells into cardiomyocytes using a previously published protocol (Supplementary Material, Fig. S2) (20). hiPSCs initially expressed high levels of TRA-1-60 and three key pluripotency markers, which rapidly decreased by 7 days of differentiation (20). Individual cells stained positively for cardiac troponin T (Fig. 5A). We observed variation in TRPM7 expression during differentiation, with cardiomyocytes at day 21 expressing more than double that observed in hiPSC cells, but this was not significant (Fig. 5B). In contrast, pluripotency markers rapidly decreased as the cell differentiated (Supplementary Material, Fig. S2C).

Beating hiPSC-CMs were seeded onto gelatin-coated glass coverslips at low density at days 15–16 for single-cell patch-clamp analysis to detect TRPM7 current at days 21–23. We found in all contractile cells subjected to whole-cell patch-clamp, typical TRPM7 currents, which were detected after dialysis of cells with Mg2+ chelating intracellular solutions. Although initial break-in currents were minimal, outward currents at +80 mV were detected after 5–10 min (Fig. 5C, E and F). We perfused the bath solution with magnesium to confirm these currents were generated by TRPM7 (Fig. 5D–F). We found that at both +80 and –80 mV, these currents were susceptible to magnesium inhibition (P < 0.001), which could be reversed following washout with standard extracellular solution.

Mimicking TRPM7 reduction using siRNA does not adversely affect action potential parameters in hiPSC-derived cardiomyocytes

To mimic possible haploinsufficiency of TRPM7 in developing cardiac cells due to p.G179V and p.T860M mutants, we used siRNA to reduce mRNA levels. To patch-clamp cells that had taken up TRPM7 siRNA, we co-transfected siGLO. Whole-cell patch clamp analysis showed a markedly reduced outward currents at +80 mV (P = 0.001, Fig. 6A and B) in cells transfected with TRPM7 siRNA. On average, this led to a 25.5 pa/pF reduction in current density, a similar reduction to that seen in 10 mM Mg2+ treated cells (28.1 pa/pF). Perhaps due to their small initial size (<5 pa/pF), inward currents at –80 mV were not significantly different between control cells, those perfused with 10 mM Mg2+ or those transfected with TRPM7 siRNA (Fig. 6C). RT-qPCR analysis of these cells found a significant decrease in TRPM7 mRNA levels in siRNA-transfected populations (Supplementary Material, Fig. S3). After confirming TRPM7 knockdown, we recorded triggered action potentials in a separate set of experiments (Fig. 6D and E). To establish whether there were any deleterious electrophysiological effects caused by reducing TRPM7 ion channel expression, we analyzed action potential morphology (Fig. 6C–L). We found that there were no statistical differences in action potential duration, minimum diastolic potential, resting membrane potential and peak voltage amplitude in the action potentials of cells transfected with scrambled or TRPM7-targeted siRNA and untransfected cardiomyocytes.

Discussion

Almost half of stillbirths go unexplained, despite extensive post-mortem analysis. In this study, we present data from previously sequenced and predicted damaging TRPM7 genetic variants found in unexplained stillbirth cases (5). There is extensive research studying ion channelopathies that cause sudden cardiac death in young adults; however, there is little evidence of harmful ion channel variants in stillbirth (21). TRPM7 is required for mammalian cardiac development, and genetic variation within the gene has been linked to a prolonged QT interval (7,15). To investigate the role these variants may play in perturbing the function of the TRPM7 ion channel, we used heterogeneous expression and hiPSC-CMs to study channel function, magnesium response and expression.

Here, we show that three TRPM7 variants modify ion channel function in comparison to the WT channel and identify TRPM7 as an ion channel present in hiPSC-derived cardiomyocytes. First, we expressed and characterized the WT channel in HEK293 and CHO-K1 cells before analyzing the effects of the four TRPM7 mutants. We report that p.G179V and p.T860M variants reduced outward currents significantly compared to the WT channel. Conversely, in a CHO-K1 specific manner, p.R494Q increased observed current density at both +80 and –80 mV. This increase remained significantly different despite inhibition with 1 mM Mg2+.

Protein expression data confirmed that decreased current density caused by the p.G179V and p.T860M variants was due to reduction in protein expression. In the case of p.T860M, this was at least in part due to rapid proteasomal degradation. It has been shown that TRPM7 is required for proper embryonic development and cardiogenesis in mice (15,16). Targeted cre-mediated knockdown of TRPM7 before E10 leads to congestive heart failure due to lack of myocardial compaction. Later deletion at E12.5 of gestation can lead to cardiomyopathy, heart
block and impaired repolarization. Given the experimental data, it is feasible to postulate that a reduction in TRPM7 function/-
expression during gestation could have an adverse effect on
human cardiac development. Mechanistically, it is hard to
determine the mechanisms that underline this effect as TRPM7’s role in
magnesium and zinc homeostasis, alongside control of apoptosis and gene expression is still under investigation (10,12,13,22). Reduced TRPM7 expression may limit the cell’s ability to take-
up divalent cations or hinder their expulsion—alongside adverse
to a cell’s gene expression profile.

Recent work with cryo-EM microscopy has revealed the
closed state structure of mouse TRPM7 at the Ångstrom scale (23). These data show several interactions between the N-
terminal domain and the inner plasma membrane, TRP domain
or C-terminal domain. Non-synonymous variants found within
these structural motifs may destabilize the protein, resulting in
misfolding and degradation.

Using a hiPSC-derived model, we demonstrate that TRPM7
can be found in immature human cardiomyocytes 21 days
post differentiation. Analysis of temporal TRPM7 expression
during cardiomyocyte differentiation showed a gradual increase
in mRNA, alongside the reduction in pluripotency factors. Single-cell patch-clamp revealed that these cells possess
relatively large quantities of TRPM7 present at the plasma
membrane, with mean outward currents of (~33.3 pA/pF). Both
treatments with Mg2+ and siRNA transfection significantly
reduced these currents, strongly suggesting their identity as
TRPM7. There is evidence to suggest that TRPM7 can regulate
expression during hiPSC-derived cardiomyocyte maturation. It is notable that increases in TRPM7 expression occur relatively early in hiPSC cardiac differentiation. It is an open issue whether TRPM7 currents contribute significantly to the fetal cardiac action potential. Our experiments with siRNA suggest that this isn’t the case under physiological ionic conditions, particularly due to the inhibitory effect of intracellular Mg2+ present in the tissue. Alternatively, haploinsufficiency may have deleterious effects that are not solely related to effects on cardiac excitability but to effects on cell survival mediated by the kinase domain.

In summary, we found two variants in TRPM7, p.G179V and p.T860M, reduce ion channel current expression, which in the case of p.T860M is likely due to rapid degradation mediated by the proteasome. In addition, the p.R494Q TRPM7 variant significantly increases TRPM7 ion channel current, in a cell-type specific manner. For the first time, we report that a TRPM7 current is present in hiPSC-CMs. However, siRNA inhibition at day 15–16, despite reducing TRPM7 current, did not alter AP duration. We believe that TRPM7 may play a key role in ensuring correct cardiac development of the fetus. TRPM7 is the major Ca2+ channel in atrial fibroblasts, and up-regulation of these transients is reported in patients with atrial fibrillation (24). Significant increases in these Ca2+ influxes from p.R494Q TRPM7 in the developing atrial myocardium could predispose developing fetal hearts to atrial fibrillation. In a similar fashion, a reduction in TRPM7 current may also predispose to cardiac arrhythmia. For example, it is known that both long and short QT syndromes arising from loss and gain of function mutations, respectively, in KCNQ1 can result in sudden cardiac death in afflicted families (25). Therefore, any changes to TRPM7 abundance during early cardiogenesis may have severe downstream effects later in gestation, potentially creating a pro-arrhythmic cardiac environment. Linking causation with genetic variation is inherently difficult, but functional assays of potential impairment of protein function can contribute significantly to assessing potential pathogenicity (26). Our data indicate that inherited or de novo TRPM7 mutations may increase the risk of unexplained stillbirth and this warrants further attention.

Materials and Methods

Cell culture, plasmids and transfection

Human Embryonic Kidney-293 cells (HEK293) were a gift from Professor Lily Jan (Howard Hughes Medical Institute, San Francisco, USA). Chinese Hamster Ovary-K1 (CHO-K1) cells were obtained from the European Collection of Authenticated Cell Cultures (supplied by Sigma—85051005). HEK293 and CHO-K1 cells were cultured in minimum essential medium (31095029, Gibco, ThermoFisher) and Ham’s F12 medium (N6658, Sigma-Aldrich), respectively. Both media were supplemented with 10% FBS (10500056, Gibco, ThermoFisher) and 100 units/ml of penicillin and 100 μg/ml of streptomycin (15140122, Gibco, ThermoFisher).

A pcDNA4/TO plasmid containing the TRPM7 gene (RefSeq NG_021363.2, NM_017672.6), and a tetracycline repressor plasmid, pcDNA6/Tet Repressor were both kind gifts from Professor Schmitz (11). An eGFP plasmid (Clontech) was used to identify successfully transfected cells. Transfection was carried out using FuGENE® HD (E2311, Promega) or NovaCHOice® (72622-3, Merck) as per manufacturer’s instructions. Briefly, reactions of TRPM7 vector and Tet Repressor plasmid were mixed with the transfection reagent in 37°C Opti-MEM® reduced serum media.
Figure 5. A TRPM7-like current can be found in 21-day-old hiPSC-derived cardiomyocytes. (A) Staining of hiPSC-CMs after 21 days of differentiation for cardiac troponin (cTnT) and DAPI. (B) qPCR analysis of TRPM7 expression levels over the course of hiPSC-CM differentiation. TRPM7 expression at each time point was normalized to GAPDH expression before being compared to TRPM7 expression in undifferentiated hiPSC cells. (C) A representative TRPM7 trace from whole-cell hiPSC-CMs. Initial current is shown (black) at initial current recordings before magnesium is removed from the bath, and a steady plateau is reached (red). (D) Time course recording of 22 day post-differentiation hiPSC-CM showing TRPM7 whole-cell current run-up and transient inhibition of 10 mM Mg2+ at +80 mV (red) and −80 mV (black). (E) Paired individual cell current measurements at +80 mV peak during extracellular solution perfusion, cells were then treated with 10 mM Mg2+ over several minutes until current plateaued. (F) Inward current recorded at −80 mV in the same cells analyzed in (D), after current stabilization and after 10 mM Mg2+ treatment. * = P < 0.05, ** = P < 0.001.

After 5–6 h, transfection complexes were removed and fresh media added. Precisely 24 h later, growth media was changed, this time supplemented with 1 μg/ml of tetracycline (87128, Sigma-Aldrich). Cells were harvested/analyzed 24–48 h after tetracycline induction.

Patch-clamp electrophysiology

Whole-cell patch-clamp recordings were made at room temperature (21–25°C), using a multiclamp 700B and digitized using a Digidata 1500B (Molecular Devices). The internal pipette contained (in mM) 145-Cs-methanesulfonate (CsSO₃CH₃), 8 NaCl, 10 HEPES and 10 EGTA; pH 7.2 with CsOH. The extracellular solution contained (in mM) 135 NaCl, 5.4 CsCl, 10 HEPES, 10 glucose, 0.1 CdCl₂ and 1 CaCl₂ (pH 7.4 with NaOH). For measuring TRPM7 current in hiPSC-CMs, the internal pipette solution contained (in mM) 120 L-aspartic acid, 20 CsCl, 2.5 EGTA, 2.5 EDTA, 10 HEPES, 120 CsOH and 5 Na₂GTP (pH 7.2 with CsOH). Extracellular solution contained (in mM) 135 NaCl, 5.4 CsCl, 10 HEPES, 10 glucose, 0.1 CdCl₂ and 1 CaCl₂ (pH 7.4 with NaOH). To record action potentials, internal pipette solution contained (in mM) 110 K-gluconate, 20 KCl, 10 HEPES, 0.05 EGTA, 0.5 MgCl₂, 5 Mg ATP, 0.3 Na₂-GTP and 5 Na₂ phosphocreatine (pH 7.4 with KOH). Extracellular solution for action potential recordings (mm): 140 NaCl, 2.7 KCl, 1 MgCl₂, 2 CaCl₂, 0.5 Na₂HPO₄ and 5 glucose (pH 7.4 with NaOH). Only cells with membrane resistance >800 MΩ and access resistance <10 MΩ were used in analysis.

Pipettes were backfilled with intracellular solution to resistance 2–3 MΩ. Pipette tips were coated with SigmaCote to reduce pipette capacitance (SL2, Sigma-Aldrich). Series resistance was
Figure 6. siRNA-mediated knockdown of TRPM7 at day 16 of differentiation does not alter hiPSC-CM AP parameters. (A) Comparison of representative TRPM7 currents after run-up in hiPSC-CMs with (black) or without TRPM7 siRNA transfection (red). (B) TRPM7 current measured at +80 mV following current run-up in non-targeting siRNA-treated hiPSC-CMs and the effect of magnesium addition or transfection with TRPM7 siRNA. (C) Inward –80 mV current measured from the same cells as in (B). (D) Representative action potential trace of non-targeting siRNA-treated hiPSC-CMs and the effect of magnesium addition or transfection with TRPM7 siRNA. (E) Representative action potential trace of hiPSC-CM transfected with siRNA-TRPM7. (F) Representative action potential trace of hiPSC-CM transfected with siRNA-scrb. (G) Maximal diastolic potential. (H) Resting membrane potential. (I) Peak voltage following action potential triggering. (J) Time to return to RMP or full action potential duration in hiPSC CMs. (K and L), APD50 and APD90 measurements from the same cells in (J). All recordings were done in 20- to 23-day-old hiPSC-CMs at room temperature. Scr = Scrambled. *** = P < 0.001 vs control.

set to 70% compensation using the in-built amplifier circuitry. After whole-cell configuration was established, recording began immediately to observe current run-up. We used a ramp voltage protocol to measure TRPM7 current: the holding potential was set at 0 mV before lowering to –100 mV and then raised to +100 mV over 250 ms before returning to 0 mV. For time-course analysis experiments, this was measured every 5 s from electrical access being established through the cell. Data were analyzed using Clampfit 10.5 (Molecular Devices).

In hiPSC-derived cardiomyocytes (hiPSC-CMs), we only analyzed triggered action potentials (APs) from cells with spontaneous action potentials. APs were triggered using a current pulse (between 400 and 800 pA) at a frequency of 1 Hz. To calculate action potential duration, the time (in ms) was
calculated as the time it took for membrane potential to return to 50 and 90% (APD50 and APD90) of the resting membrane potential after current trigger. Cells were paced for 50 s before recording the final 10 action potentials. These results of the final 10 traces were combined to give an ‘average’ action potential for every cell. We did not analyze cells according to ventricular-like or atrial-like categorizations due to a limited number of cells.

Protein harvesting, SDS-PAGE and western blotting
HEK293 cells were transfected at ~70% confluence as previously mentioned and harvested 24 h after tetracycline induction. Cells were washed with PBS+ (0.1 mM CaCl2 and 1 mM MgCl2) before a 30-min incubation on ice with NP-40 lysis buffer containing protease inhibitors (1181617001, Sigma-Aldrich). Lysates were scraped into pre-chilled tubes and cleared by centrifugation at 10,000g for 10 min at 4°C. Amount of 20 µg of protein was loaded for each well into 6% polyacrylamide gels. Proteins were transferred to PVDF membranes overnight and then blocked for 1 h in PBS-T with 5% milk powder or 1% BSA (A2153, Sigma-Aldrich). Membranes were then incubated with either anti-TRPM7 (sc-271099, Santa Cruz) or β-actin (ab8226, Abcam) antibodies for 1 h at room temperature. Secondary antibodies were incubated for 1 h at room temperature before imaging with Amersham ECL western blotting detection reagent (GE Healthcare). Primary and secondary antibody details are listed in Supplementary Material, Table S2. ImageJ software was used to quantify TRPM7 expression, which was normalized to loading control β-actin levels, and then all stillbirth variant expression was normalized to the WT value in the same gel. For proteasomal inhibition experiments, cells were treated with 5 µM MG132 (Sigma-Aldrich) for 16 h prior to harvesting.

RNA harvesting and quantitative PCR
Total cell RNA was harvested using a RNeasy mini kit (Qiagen) as per manufacturer’s instructions either 24 h after tetracycline induction or at specific time points of stem cell differentiation. During harvesting of RNA, the column was treated with DNase I for 15 min at room temperature. RNA was purified in nuclease-free water, and 50 ng used per qPCR reaction. All probes used are listed in Supplementary Material, Table S3. Data were analyzed using 2^(-ΔΔCt) method normalized to loading control and then normalized to WT. To calculate gene expression in hiPSC-derived cardiomyocytes, RNA levels were first normalized to the housekeeping gene GAPDH. All expression values were then normalized to the expression of the gene of interest in naïve iPSC cells prior to differentiation.

Confocal microscopy
Cells were first washed with PBS+ to remove media and then fixed with 3.7% formaldehyde (Sigma-Aldrich) for 10 min at room temperature. Cells were permeabilized with PBS Triton X-100 0.1% for 10 min. We blocked cells in PBS-T 5% milk for 1 h and carried out primary anti-cTnT incubation for overnight at 4°C. Fluorescent secondary antibody incubation took 1 h at room temperature. All antibody details are listed in Supplementary Material, Table S1. Nuclei were counter stained with DAPI and all slides stored at 4°C until imaging with an LSM710 confocal microscope (Zeiss).

Human stem cell-derived cardiomyocytes
Healthy human-induced pluripotent stem cell line (named—HS1M) was generated and maintained as previously described (27). Cardiomyocytes were differentiated using a protocol based upon previous work published by Burridge et al. (20). In brief, hiPSCs were seeded at 150,000 cells/cm² in a 24-well plate and allowed to reach ~80% confluence over 48 h. Media was changed to ‘CDM3’ (RPMI1640 (Thermofisher) supplemented with 500 µg/ml O. sativa-derived recombinant human albumin (Sigma-Aldrich), 213 µg/ml L-ascorbic acid 2-phosphate (Sigma-Aldrich)) with 6 µM CHIR99021 (Sigma-Aldrich). After 48 h, media was changed to CDM3 supplemented with 2 µM C59 for a further 2 days before regular CDM3 media changes every 2–3 days. For patch-clamp analysis, at 15 days post-differentiation, cells were treated with TrypLE Express (Thermofisher) for 10 min at 37°C and seeded onto glass coverslips at a density of 10,000 cells/cm².

For TRPM7 siRNA knockdown experiments, 48 h before patch-clamp analysis, adhered cells were transfected with either SMARTpool: ON-TARGETplus TRPM7 targeting siRNA or a scrambled control (L-00593-00-0005 or L-006233-00-0005, respectively, Thermofisher). DharmaFECT transfection reagent 1, 2.5 µl/well (T-2001-02, Thermofisher) was used to transfect 2.5 µl/well of 2 µM siRNA.

Statistics
All data were analyzed with GraphPad Prism 6. Grouped data were tested with one-way ANOVA, using Dunnett’s multiple comparison to test variants vs WT unless otherwise stated. For individual analysis of two groups, unpaired t-tests were used to calculate statistical differences. When comparing two independent variables, two-way ANOVA with Sidak’s multiple comparison test was used. All group data are presented as mean ± S.E.M. Significant P values are denoted as * = P < 0.05, ** = P < 0.01 and *** = P < 0.001.

Supplementary Material
Supplementary Material is available at HMG online.

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References


