In silico investigation of a KCNQ1 mutation associated with short QT syndrome

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Short QT syndrome (SQTS) is a rare condition characterized by abnormally 'short' QT intervals on the ECG and increased susceptibility to cardiac arrhythmias and sudden death. This simulation study investigated arrhythmia dynamics in multi-scale human ventricle models associated with the SQTS-related V307L KCNQ1 'gain-of-function' mutation, which increases slow-delayed rectifier potassium current (I\textsubscript{Ks}). A Markov chain (MC) model recapitulating wild type (WT) and V307L mutant I\textsubscript{Ks} kinetics was incorporated into a model of the human ventricular action potential (AP) for investigation of QT interval changes and arrhythmia substrates. In addition, the degree of simulated I\textsubscript{Ks} inhibition necessary to normalize the QT interval and terminate re-entry in SQTS conditions was quantified. The developed MC model accurately reproduced AP shortening and reduced effective refractory period associated with altered I\textsubscript{Ks} kinetics in homozygous (V307L) and heterozygous (WT-V307L) mutation conditions, which increased the lifespan and dominant frequency of re-entry in 3D human ventricle models. I\textsubscript{Ks} reductions of 58% and 65% were sufficient to terminate re-entry in WT-V307L and V307L conditions, respectively. This study further substantiates a causal link between the V307L KCNQ1 mutation and pro-arrhythmia in human ventricles, and establishes partial inhibition of I\textsubscript{Ks} as a potential anti-arrhythmic strategy in SQTS.

The short QT syndrome (SQTS) was first published as a distinct clinical entity in 2000\textsuperscript{1}. It is characterised by an abnormally short QT interval on the electrocardiogram (ECG), tall and peaked T-waves, poor rate adaptation of the QT interval, shortened atrial and ventricular refractory periods, increased risk of atrial and ventricular arrhythmias, and an increased incidence of sudden death in affected patients\textsuperscript{2–7}. To date, six genetic variants of the SQTS have been identified (SQT1–6) with the first three variants (SQT1–3) being caused by gain-of-function mutations to genes encoding different K\textsuperscript{+} channel subunits while SQT4–SQT6 are due to loss-of-function mutations to genes encoding subunits responsible for L-type calcium channel current (I\textsubscript{Ca_L}), resulting in a mixed SQT-Brugada phenotype\textsuperscript{8,9}. SQT2 is caused by mutations to KCNQ1 (KvLQT1)\textsuperscript{10–13}, which in combination with KCNE1 encodes the proteins responsible for human cardiac IKs channel\textsuperscript{14,15}. The first-identified adult SQT2 variant involves a G \textrightarrow{} C substitution in nucleotide 919 of KCNQ1 that leads to an amino acid substitution of valine to leucine on residue 307 (V307L) in the KCNQ1 channel pore helix\textsuperscript{10}. The V307L mutation shifts the voltage dependence of KCNQ1+KCNE1 activation towards more negative voltages and accelerates channel activation, which consequently leads to increased current during ventricular repolarisation\textsuperscript{10}. For the V307L KCNQ1 SQT2 mutation, initial in silico data\textsuperscript{10} were able to reproduce action potential (AP) shortening but did not address directly effects on the QT interval or arrhythmia mechanisms. In a subsequent study\textsuperscript{16}, we developed a Hodgkin-Huxley (HH) style model for the V307L KCNQ1 mutation and investigated its effects in idealised one-dimensional (1D) and two-dimensional (2D) tissue. However, that study\textsuperscript{16} had some intrinsic limitations: (i) the developed HH model of SQT2 did not incorporate slow deactivation of the I\textsubscript{Ks} with the V307L KCNQ1 mutation that was subsequently identified\textsuperscript{17}; (ii) it did not consider possible functional

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consequences of realistic ventricular anatomical geometry in two and three dimensions, which play important roles in initiation and maintenance of arrhythmic excitation waves18, 19.

One aim of this study was to address the above limitations of the previous study using a HH model of SQT216. We conducted the present study in order to: (i) develop a novel biophysically-accurate and validated Markov chain (MC) model to recapitulate the kinetic changes to \( I_{Ks} \) in the SQT2 V307L KCNQ1 mutation based on available experimental data at physiological temperature; (ii) determine the functional consequences of the SQT2 V307L mutation on AP repolarisation and the QT interval by incorporating it into a well-established human ventricular cell model20; (iii) explore the arrhythmogenic substrate in the SQT2 V307L mutation by using “realistic” 2D tissue and 3D organ-scale simulations; and, moreover; (iv) investigate theoretically the degree of \( I_{Ks} \) inhibition required to normalise the QT interval as a pseudo-pharmacological therapeutic intervention.

The multi-scale cardiac modelling approach taken in this study has been employed successfully in our previous studies to dissect ionic mechanisms underlying QT interval shortening and pro-arrhythmia in variants 1 and 3 of the SQT318, 19. The developed MC model represents a significant advance over previous studies10, 18 in SQT2 modelling and the reported results further understanding of the mechanisms by which the SQT2 V307L mutation enhances susceptibility to reentrant arrhythmia. Furthermore, a theoretical basis for pharmacological intervention in SQT2 is proposed.

Results

Simulation of single cell \( I_{Ks} \) under control and V307L mutation conditions. First, we tested the ability of the \( I_{Ks} \) MC model to reproduce the previously published experimental data2 on the voltage dependence of wild type (WT) and V307L KCNQ1 + KCNE1 \( I_{Ks} \) at physiological temperature. Figures 1Aii and 1Bii show the voltage clamp protocol used which is the same as that used for the experimental data2 and the generated \( I_{Ks} \) current traces (Fig. 1Aii and Bi) from which the I-V relationships were reconstructed (Fig. 1Aiii and Bi). The simulated I-V relationships for both the WT and KCNQ1 V307L conditions match those recorded experimentally17. The simulated current traces match experimental recordings; under the V307L mutation condition, the slower deactivation rate and faster activation rate of the \( I_{Ks} \) channel21 compared to the WT condition were reproduced (Fig. 1Aii and Bi).

To validate the MC model formulations, we next simulated AP voltage clamps to test their ability to reproduce the dynamic properties of WT and V307L KCNQ1 mutant \( I_{Ks} \) channels. Figure 1Aiv and Biv show the results of the simulated AP clamp experiments compared to those obtained in vitro17. The MC model reproduced the profile of \( I_{Ks} \) during imposed AP commands with a high degree of accuracy, both in the WT condition and with augmented \( I_{Ks} \) in simulations incorporating the KCNQ1 V307L mutation.

Following incorporation of the WT and V307L MC models into the 2006 model of ten Tusscher et al. (TNNP)26, Fig. 2 shows simulated APs (Ai), \( I_{Ks} \) profile (Aii) and \( I_{Ks} \), instantaneous I-V relationship for an epicardial (EPI) cell (Ai). The mid-myocardial (MIDDLE) and endocardial (ENDO) counterparts are shown in Fig. 2B and C, respectively. WT \( I_{Ks} \), increased progressively following the upstroke of the AP and reached maximal amplitude very late during the plateau phase before declining rapidly during terminal repolarisation. WT-V307L \( I_{Ks} \), activated earlier than WT and increased in amplitude more rapidly. Compared to WT current, it reached significantly higher maximal amplitude early during the plateau leading to abbreviation of the AP duration (APD). V307L \( I_{Ks} \), activated even earlier than WT-V307L \( I_{Ks} \), increased even more rapidly, and attained higher maximal amplitude. Consequently, it abbreviated the APD to a greater extent. Under the WT condition, the computed APD\(_{90}\) was 326 ms, 454 ms and 327 ms for EPI, MIDDLE and ENDO cells respectively. These were shortened respectively to 233 ms, 355 ms and 234 ms under the WT-V307L condition, and to 194 ms, 306 ms and 194 ms under the V307L mutation condition. The APD\(_{90}\) values for all the conditions are summarised in Table 1. The APD shortening resulted from the augmented \( I_{Ks} \) early during the plateau phase of the AP as shown by the time course of \( I_{Ks} \) (Fig. 2Aii–Cii) and the I-V phase plots in Fig. 2Aii–Cii. The greatest APD shortening was observed for the MIDDLE cell model (Table 1).

The APD abbreviation was rate-dependent as shown by the APD restitution (APD-R) curves in Fig. 3A–C for the EPI, MIDDLE and ENDO cell types, respectively. Over the range of diastolic intervals (DI) studied, the APD was smaller in the WT-V307L and V307L mutant conditions than in the WT condition. The mutations also steepened the APD-R curves in each cell type as shown by the computed maximal slopes for each APD-R curve in Fig. 3D. In the EPI cell, the maximal slopes of the WT-V307L and V307L conditions were similar, while there was a progressive increase in steepness of the slopes in the MIDDLE cell type. In the ENDO cell, while the slopes under the WT-V307L and V307L conditions were steeper than WT, the slope of the WT-V307L condition was steeper than that of the V307L mutant. The ERP reduction was also rate-dependent. It was reduced under the WT-V307L and V307L mutation conditions compared to the WT condition across the range of basic stimulus cycle lengths (BCL) as shown in ERP-R curves in Fig. 3E–G for the EPI, MIDDLE and ENDO cell types, respectively. In the EPI and ENDO cells, there was little difference in the slopes of the ERP-R curves between the WT, WT-V307L and V307L conditions (Fig. 3H). The slope was steeper for the mutation conditions in the MIDDLE cell compared to WT but the slope for the WT-V307L condition was slightly steeper than that for the V307L mutant alone. The mutations also shifted the ERP-R curve lower-right implying that the KCNQ1 V307L mutation enabled ventricular cells to support higher rate electrical activity (as normally seen during VT and VF conditions).

Simulation of the ECG with WT and SQT2 mutant \( I_{Ks} \). Using a 1D strand model of the ventricular wall, we computed a pseudo-ECG under the WT, WT-V307L and V307L KCNQ1 conditions (Fig. 4D–F). These were extracted from a propagating wave from the ENDO end of the strand towards the EPI end (Fig. 4A–C). Time runs horizontally from left to right in Fig. 4A–C while space runs vertically from the ENDO end at the bottom to the EPI end at the top. The QT interval was shortened from 351 ms in the WT condition to 292 ms in the WT-V307L.
condition and to 262 ms in the V307L condition (Fig. 4G). T wave width (measured as the time interval between $T_{\text{peak}}$ and $T_{\text{end}}$) also changed from 49 ms (WT) to 60 ms (WT-V307L) and 64 ms (V307L). These simulations thus reproduce the key features observed in the ECGs of SQTS patients; abbreviated QT interval, tall and peaked T-waves and wider $T_{\text{peak}}$ to $T_{\text{end}}$.

As the only difference between these simulations is the altered kinetics of $I_{Ks}$ resulting from the mutation, the observed changes in the QT interval, T wave height and width can be confidently attributed to the parameters corresponding to the V307L mutation.

Gima and Rudy\textsuperscript{21} showed that increased spatial gradient of membrane potential was responsible for the increase in T-wave amplitude during simulated hyperkalaemia. In order to determine if the same effect was responsible for the taller T wave amplitudes in the WT-V307L and V307L ECGs (Fig. 4D–G), we examined the effects of this KCNQ1 mutation on membrane potential heterogeneity ($\delta V$) in supplementary simulations.

Supplementary Figure S2 shows the pairwise differences between EPI, MIDDLE and ENDO cells during an AP. Under the KCNQ1 WT-V307L and V307L mutation conditions, the maximal $\delta V$ between EPI-MIDDLE and ENDO-MIDDLE cells were greater than under the WT condition, which contributed to the augmented T-wave.

Figure 1. Simulated voltage and AP clamp experiments for $I_{Ks}$. Current traces for WT (Ai) and V307L (Bi) KCNQ1 $I_{Ks}$ at step potentials of $-60$ mV (blue), $-30$ mV (green), $0$ mV (red) and $30$ mV (cyan) using the voltage clamp protocol shown in (Aii,Bii). Comparison of simulated and experimental I-V relationship for $I_{Ks}$ under WT (Aiii) and V307L (Biii) conditions. $I_{Ks}$ profile during AP clamp using a ventricular AP waveform voltage command in WT (Aiv) and V307L (Biv) conditions. All experimental data, including insets shown in panels Ai and Bi, are taken from El Harchi et al.\textsuperscript{17}.
amplitude$^{16,18,21}$. Also shown is the spatial distribution of APD$_{90}$, the spatial gradient of APD$_{90}$, and the absolute value of the spatial gradient of APD$_{90}$. The spatial gradient of APD$_{90}$ was augmented across the strand but markedly so in the ENDO region.

**Investigating the arrhythmogenic substrate in SQT2 – 1D simulations.** In further supplementary simulations using the 1D strand, we investigated the vulnerability of WT, WT-V307L and V307L tissue to unidirectional block in response to a premature stimulus applied during the refractory tail of a previous excitation wave. During such conduction block tissue is rendered more susceptible to re-entrant arrhythmia. Supplementary Figure S3 shows the percentage increase compared to the WT condition in the temporal vulnerable window to conduction block, measured at a location 5.0 mm away from the epicardial end of the 1D strand. The width of the vulnerable window was increased by 37% and 82% with respect to the WT in WT-V307L and V307L conditions, respectively.

**Investigating the arrhythmogenic substrate in SQT2 – 2D and 3D simulations with realistic geometry.** In a realistic human ventricle cross-sectional slice (Fig. 5), we investigated the response of WT, WT-V307L and V307L tissue to a local premature stimulus applied within the left ventricular wall during the tissue’s vulnerable window (WT: 370 ms after the arrival of conditional wavefront; WT-V307L: 310 ms after the arrival of conditional wavefront; V307L: 230 ms after the arrival of conditional wavefront). The results of 2D simulations are shown in Fig. 5 (and Supplementary Videos S1–3). Following the premature stimulus, a re-entrant excitation wave was initiated within the left ventricular wall as shown in Fig. 5A–D for WT, Fig. 5Aii–Dii for WT-V307L and Fig. 5Aiii–Diii for the V307L condition. The snapshots shown in Fig. 5A–D show subsequent conduction of the induced re-entrant excitation wave from the applied premature stimulus for the WT (Fig. 5B–D), WT-V307L (Fig. 5Bii–Dii) and V307L (Fig. 5Aiii–Diii) conditions. Under the WT condition, the initiated

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**Table 1.** Action potential duration in WT, WT-V307L and V307L conditions. Action potential duration (APD) was measured in all transmural cell types; epicardial (EPI), mid-myocardial (MIDDLE), and endocardial (ENDO) at a frequency of 1 Hz. ΔAPD is measured relative to WT.

|       | WT (ms) | WT-V307L (ms) | V307L (ms) |
|-------|---------|---------------|------------|
| EPI   | 325.6   | 233.0         | 193.6      |
|       |         | ΔAPD 92.6     | ΔAPD 132.0 |
| MIDDLE| 453.6   | 355.4         | 305.9      |
|       |         | ΔAPD 98.2     | ΔAPD 147.7 |
| ENDO  | 327.4   | 233.8         | 194.0      |
|       |         | ΔAPD 93.6     | ΔAPD 133.3 |

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**Figure 2.** Action potentials and $I_{Kr}$ profiles. Steady state APs (i) for EPI (A), MIDDLE (B), and ENDO (C) cells in WT (blue), WT-V307L (green), and V307L (red) conditions at a pacing rate of 1 Hz. Corresponding $I_{Kr}$ profiles (ii) and I-V relationships (iii) for EPI (A), MIDDLE (B), and ENDO (C) cells.
re-entry self-terminated after 1.1 s (Fig. 5Di, Supplementary Video S1) but it persisted under the mutation conditions throughout the whole simulation period (5 s) (WT-V307L: Fig. 5Bii–Dii, Supplementary Video S2; V307L: Fig. 5Biii–Diii, Supplementary Video S3).

The time course of an AP in the left ventricle is shown for the WT, WT-V307L and V307L conditions in Fig. 5Ei–Eiii respectively. Power spectrum analysis of the recorded whole-field averaged electrical activity from the tissue revealed a higher dominant frequency in the mutation conditions (3.32 Hz for WT-V307L and 4.30 Hz for V307L compared to the WT condition (1.96 Hz) (Fig. 5F). These 2D simulation results illustrate that the KCNQ1 V307L mutation increases tissue susceptibility to arrhythmogenesis by maintaining re-entrant excitation waves.

Realistically, the ventricles are three-dimensional and have a much more complex anisotropic geometry compared to the 2D ventricular slice. Therefore, one cannot necessarily assume that sustained reentry in the 2D tissue model translates to similar activity in 3D tissue. Consequently, we performed further simulations with a 3D anatomical human ventricle geometry. The results are shown in Fig. 6, which shows snapshots of the evolution of re-entrant scroll waves (WT: Fig. 6Ai–Di, Supplementary Videos S4 and S5; WT-V307L: Fig. 6Aii–Dii, Supplementary Videos S6 and S7; V307L: Fig. 6Aiii–Diii, Supplementary Videos S8 and S9) developing as a response to a premature stimulus. For the WT condition, the scroll wave self-terminated with a lifespan of 0.5 s (Fig. 6F). However, under WT-V307L and V307L mutation conditions, the scroll wave broke up forming multiple re-entrant wavelets that self-terminated within 2.5 s in WT-V307L tissue but were sustained throughout the 5 s simulation period in V307L tissue (Fig. 6F). Power spectrum analysis of the registered pseudo-ECG shows the dominant frequency of ventricle excitation to be 2.34 Hz for the WT condition, 3.13 Hz for the WT-V307L mutation condition and 7.42 Hz for the V307L mutation condition (Fig. 6G). Figure 6Ei–Eiii show a recording of the evolution of the AP of a cell in the left ventricle for the WT, WT-V307L and V307L conditions.

Investigating \( I_{Ks} \) as a potential therapeutic target in SQT2. As a pseudo-pharmacological approach to treating patients with SQT2, we mimicked \( I_{Ks} \) channel blockade by drugs in order to determine the extent of blockade required to normalise the QT interval. To investigate this, maximal channel conductance of mutant \( I_{Ks} \) was reduced in the EPI, MIDDLE and ENDO single cell types. Figure 7A–E show the results for an EPI cell in the TNNP model under the WT-V307L (Fig. 7A–B) and V307L (Fig. 7D–E) conditions. In all three cell types, approximately 60% simulated \( I_{Ks} \) blockade was required to make the APD comparable to that of WT under the
WT-V307L mutation condition while approximately 76% $I_{Ks}$ blockade was necessary under the V307L mutation condition.

Using the intact 1D strand, we performed a similar investigation on the normalization of the QT interval. In tissue, due to the electrical coupling between cells via gap junctions, APD differences between different cell types are reduced. Thus, for the WT-V307L mutation condition, 59% $I_{Ks}$ reduction (Fig. 7C) was needed to normalise the QT interval to that of WT while under the V307L mutation condition, approximately 75% $I_{Ks}$ reduction was necessary (Fig. 7F). These results are similar to the single cell situation.
Finally, we investigated whether simulated $I_{Ks}$ blockade could terminate re-entrant activity under the mutation conditions (Fig. 8) in the 3D anatomical human ventricles. Figure 8A shows the WT condition with a premature S2 stimulus applied during the tissue's vulnerable window at 355 ms. If $I_{Ks}$ reduction under the mutation conditions normalises the QT interval, then the application of a premature stimulus at this same time (355 ms) should produce somewhat similar activity to WT. Figure 8Bi and Ci show WT-V307L and V307L mutant tissue with no $I_{Ks}$ reduction, i.e. the pure heterozygote and homozygote mutant conditions respectively with a premature stimulus applied at 315 ms for WT-V307L and 260 ms for V307L leading to re-entrant activity that persists beyond the...
WT reentry lifespan. It transpired that 58% I_Ks reduction was sufficient to make the WT-V307L reentry lifespan (Fig. 8Bii) comparable to that of WT while 65% I_Ks reduction was adequate under the V307L mutation condition (Fig. 8Cii). These simulations illustrate the possibility of I_Ks as a relevant drug target to treat tachyarrhythmia in the SQT2 setting.

Discussion
In this study, we have developed a MC model of I_Ks incorporating the SQT2 V307L KCNQ1 mutation in order to elucidate possible pro-arrhythmic effects of this KCNQ1 gene mutation, and investigate in silico the possibility of I_Ks inhibition as a treatment for SQT2. The major findings of the present study are summarised as follows: (i) a novel biophysically-detailed MC model formulation that reproduces accurately the dynamic properties of the KCNQ1 V307L mutation; (ii) the V307L mutation abbreviates the APD and steepens the APD restitution.
The V307L mutation shortens the QT interval, increases T wave amplitude and T peak−Tend duration, all of which are concordant with clinical observations regarding the SQTS; significantly reduced ERP in all transmural cell types in the KCNQ1 V307L mutation accelerates re-entrant excitation waves in human ventricles; augmented membrane potential (δV) differences and transmural APD dispersion associated with the KCNQ1 V307L mutation contributes to increased T-wave amplitude and increased temporal vulnerability to uni-directional conduction block by a premature excitation at some localised regions of the ventricles; IKs blockade is a potential therapeutic means of normalizing the QT interval and terminating re-entrant activity in the SQT2 setting. These results provide increased insight into understanding the causal link between the KCNQ1 V307L mutation and QT interval shortening and tachyarrhythmia.

In the first report of the KCNQ1 V307L mutation in a patient with the SQTS, Bellocq et al.10 used a Pribe-Beuckelmann ventricular cell AP model22 to demonstrate AP shortening. In our previous study16, we used the 2004 TNNP human ventricular AP cell model23 with modified HH IKs formulations reproducing V307L KCNQ1+KCNE1 kinetics available at that time to also demonstrate AP abbreviation, QT interval shortening, T wave morphology changes, reduced minimal substrate size for re-entry and reentrant activity in idealised 2D geometry. However, the current study is the first to reproduce the subsequently identified slower deactivation of V307L KCNQ1+KCNE1 IKs and to examine the mutation’s functional consequences in realistic 2D and 3D anatomical geometries. The present study is also the first to investigate the blockade of IKs as a potential pharmacological intervention for treating SQT2 patients.

Figure 7. IKs blockade in single cell and 1D simulations. Action potentials in WT-V307L (A) and V307L (D) conditions under varying degrees of IKs blockade. The dashed line represents the WT and the boxed percentage represents the degree of IKs block required to normalize the APD under the respective mutation condition. IKs profiles corresponding to the APs shown in (A) and (D) are shown for WT-V307L (B) and V307L (E) conditions, respectively. Pseudo-ECGs corresponding to varying degrees of IKs blockade are shown in WT-V307L (C) and V307L (F) conditions. The blue line represents the WT and the boxed percentage represents the degree of IKs block required to normalize the QT interval under the respective mutation condition.
Figure 8. Termination of re-entry in 3D ventricle model by $I_{Ks}$ blockade. (A) Application of a premature S2 stimulus in a local region of WT tissue during the refractory period (355 ms) leads to the development of a scroll wave (500 ms) that terminates in under 1000 ms. (B) Application of a premature S2 stimulus in a local region of WT-V307L tissue during the refractory period (315 ms) leads to the development of a scroll wave (500 ms) which persists beyond 1000 ms. (Bii) Application of a premature S2 stimulus in a local region of WT-V307L tissue with 58% $I_{Ks}$ blockade during the refractory period (355 ms) leads to the development of a scroll wave (500 ms) that terminates in under 1000 ms. (C) Application of a premature S2 stimulus in a local region of V307L tissue during the refractory period (260 ms) leads to the development of a scroll wave (500 ms) which persists beyond 1000 ms. (Cii) Application of a premature S2 stimulus in a local region of V307L tissue with 65% $I_{Ks}$ blockade during the refractory period (355 ms) leads to the development of a scroll wave (500 ms) that terminates in under 750 ms.
The MC structure used in this study\textsuperscript{24} was previously employed in the study of O’Hara \textit{et al}.\textsuperscript{25}, in which effects of the ‘silent’ Q357R KCNQ1 mutation in the long QT syndrome (LQT1) were probed \textit{in silico}. That study\textsuperscript{28} found that multiple insults to the repolarization reserve were required to reproduce the complex arrhythmic phenotype associated with the Q357R mutation, highlighting the need for a more kinetically-accurate description of $I_{\text{Ks}}$ channel kinetics than can be achieved with traditional HH formulations. This study builds on the approach employed by O’Hara \textit{et al}.\textsuperscript{25}, incorporating a detailed MC description of $I_{\text{Ks}}$ kinetics associated with a KCNQ1 channellopathy into anatomically-detailed human ventricular tissue simulations.

The SQTS is associated with malignant tachycardias\textsuperscript{2, 4, 10, 26} and some patients present with ventricular fibrillation episodes\textsuperscript{10, 27} including the SQT2 proband\textsuperscript{10}, who was successfully resuscitated from ventricular fibrillation. In our simulations we found that the KCNQ1 V307L mutation reduces ERP at all rates (Fig. 3). This helps to maintain re-entrant activity in tissue, as it decreases the wavelength of ventricular excitation waves, allowing higher activation frequencies of re-entrant excitation waves. This is evident in the increased lifespan and dominant frequency associated with the WT-V307L and V307L mutation conditions in both 2D and 3D simulations (Figs 5 and 6). Furthermore, we found that the SQT2 mutation increases vulnerability to initiation of re-entrant activity through localized increases in transmural heterogeneity of membrane potential, which leads to an increased temporal window of vulnerability to unidirectional conduction (Supplementary Figure S3). These findings are qualitatively in accord with our previous study\textsuperscript{46} using the HH model of SQT2 with idealized tissue geometrical structures, demonstrating the elucidated arrhythmogenic mechanisms of SQT2 are model independent and further solidify the causative link between KCNQ1 V307L mutation and ventricular arrhythmogenicity.

The use of Implantable Cardioverter Defibrillators (ICDs) is the current preferred treatment for the SQTS\textsuperscript{2, 4, 28, 29}. However, as the SQTS is characterised by tall and peaked T-waves, there is the risk of inappropriate shocks to the patient due to T-wave over-sensing\textsuperscript{2, 4, 28}. Additionally, ICDs do not restore the QT interval to its normal duration and are not suitable for all patients (e.g. infants). Therefore, pharmacological alternatives that can restore the normal duration of the QT interval and offer protection from arrhythmias are being actively pursued\textsuperscript{4, 4, 30, 31}. Hydroxychloroquine has been found to be effective in prolonging the QT interval and preventing episodes of VF, but its effect is greater in patients with identified SQT1 (hERG) mutations than in those without SQT1 mutations\textsuperscript{32}. At present, there is no selective $I_{\text{Ks}}$ blocker in clinical use. \textit{In vitro} experiments have shown that although $I_{\text{Ks}}$ is selectively blocked by chromanol 293B\textsuperscript{33, 34}, the blocking potency of this agent is reduced for recombiant channels containing the V307L KCNQ1 mutation\textsuperscript{17, 34}. In contrast, $I_{\text{Ks}}$ channels incorporating the V307L KCNQ1 mutation were inhibited by the quinolone agent mefloquine with a similar potency to that for WT channels\textsuperscript{37}.

Our simulations mimicked selective pharmacological block of the $I_{\text{Ks}}$ channel in the SQT2 setting through reductions in $I_{\text{Ks}}$ magnitude. These showed that in the heterozygotic WT-V307L KCNQ1 condition, a blockade of $I_{\text{Ks}}$ by $\text{58}\%$ is sufficient to restore the QT interval to its original duration and make the tissue behave like WT tissue whereas under the V307L homozygote condition, a blockade of $I_{\text{Ks}}$ by $\text{65}\%$ is sufficient to achieve the same result. In our simulations, a high degree of $I_{\text{Ks}}$ block was required in the setting of SQT2 to ‘correct’ the cellular level APD and QT interval in tissue as well as to reduce tissue vulnerability and lifespan of reentry to those of normal tissue. This is consistent with the extent of $I_{\text{Ks}}$ augmentation arising from SQT2 conditions, however. This is demonstrated in Supplementary Figure S13. In these simulations, increasing maximal $I_{\text{Ks}}$ channel conductance in the WT condition by an amount equal to the inverse of that required to normalise the AP in SQT2 mutant conditions led to comparable shortening of the APD, and thus corresponded approximately to the amount of increased $I_{\text{Ks}}$ due to SQT2 mutant conditions. Although simulated $I_{\text{Ks}}$ block is a theoretical consideration (in the absence of clinically used selective $I_{\text{Ks}}$ inhibitors), and involves a simple current reduction rather than modification of kinetics, our study is nonetheless the first to provide proof-of-concept that $I_{\text{Ks}}$ block has the potential to be effective in the SQT2 setting. Furthermore, whilst the important role of $I_{\text{Ks}}$ in repolarization reserve means that $I_{\text{Ks}}$ blockade can lead to torsade de pointes\textsuperscript{33, 35, 36}, this risk is likely to be greatest against a background of a normal, rather than abbreviated QT interval.

Inhibition of $I_{\text{Ks}}$ as a potential therapeutic strategy was further tested in another form of SQT2 caused by the KCNQ1 V141M mutation, which differs from the V307L mutation in that it induces a constitutively active voltage-independent current component\textsuperscript{11}. Briefly, the MC was further modified to recapitulate kinetics of the V141M KCNQ1 mutation (shown in Supplementary Figure S11), including presence of a voltage-independent component, shifted voltage dependence of activation to less depolarized potentials, and significantly slowed deactivation\textsuperscript{37}. We found that the heterozygotic WT-V141M condition produced a significant shortening of the APD and QT interval, consistent with the SQTS phenotype. Simulation of a substantial degree of $I_{\text{Ks}}$ block (79%) resulted in normalization of the AP, as shown in Supplementary Figure S12. These supplementary simulations help to justify and validate our modelling approach, whilst providing further evidence that $I_{\text{Ks}}$ block may, in principle, be an effective therapeutic strategy in the setting of SQT2.

In order to determine whether or not the principal results obtained in this study pertaining to the KCNQ1 V307L mutation were model-dependent, single cell and 1D simulations performed in the TNNP model were also carried out in the more recently-developed O’Hara-Rudy dynamic (ORD) human ventricular cell model\textsuperscript{38}. At the single cell level, the major findings reported using the TNNP model were matched qualitatively by the ORd model, i.e. the SQT2 mutations significantly reduced the APD, affecting transmural regions of the ventricles differentially (Supplementary Figure S4), and increased maximum slope of AP restitution (Supplementary Figure S5). In the ORd model intact 1D strand, the MC formulation of the KCNQ1 V307L mutation accurately reproduced QT interval shortening and an increase in T-wave amplitude – two hallmarks of the SQTS (Supplementary Figure S6).\textsuperscript{2, 4–7} The underlying mechanisms contributing to increased T-wave amplitude due to the SQT2 mutations were the same as in the TNNP model, i.e. augmented maximal membrane potential difference, $\delta V$, between coupled cells in tissue (Supplementary Figure S7) due to increased spatial gradient of APD across the transmural strand (Supplementary Figure S8). Furthermore, tissue temporal vulnerability to
uni-directional conduction in the ORd model 1D transmural strand (measured at the MIDDLE-EPI border) was increased by the SQT2 mutations (Supplementary Figure S9).

$\text{I}_\alpha$ reductions of 50% and 65% in WT-V307L and V307L conditions respectively were required to normalize the APD in single cell ORd model simulations (Supplementary Figure S10). These values are comparable to the TNNP values of 60% and 76%. Similar to the single cell situation, the degrees of $\text{I}_\alpha$ blockade necessary to normalise the QT interval in SQT2 mutation conditions in the ORd model were slightly lower but still fairly close to those in the TNNP model. The model independence of these results is significant. Obtaining qualitative agreement between two distinct human ventricle cell models shows that the fundamental mechanisms by which the SQT2 mutations affect ventricular electrophysiology are the same even when there are differences in ion channel kinetics, as in the TNNP and ORd models.

Limitations of the main human ventricular cell model used (TNNP 2006) have been discussed in detail elsewhere$^{18,20,39}$. In the multicellular tissue model - due to a lack of detailed experimental data - the proportion of each region comprised each distinct cell type and intercellular electrical coupling was chosen to produce a positive T-wave and a conduction velocity of a planar solitary excitation wave close to experimental data, similar to those used in other studies$^{16,21,40}$. The heterozygote formulation used in this study relies on the simplifying assumption that V307L $\text{I}_\alpha$ behaves similar to a 50:50 mixture of WT and mutant channels. The channel population may be more complex in reality, with each channel comprising both WT and mutant KCNQ1 subunits. That said, our previous study$^{45}$ investigated the effects of varying mutant subunit composition in SQT2 based on different expression/co-expression ratios used in experiments in the original paper describing the V307L KCNQ1 mutation$^{10}$, and found that the degree of APD and QT interval shortening increased progressively with the level of V307L expression. Thus, the simplifying assumption adopted in the present study is unlikely to affect adversely the simulation predictions:

Results: our heterozygote formulation reproduced QT interval shortening and increased T wave amplitude associated with the SQT2 phenotype$^{4-47}$.

Care must be exercised in the interpretation of results from the 2D model, as it is based on a single slice of the ventricle wall. Although we considered anisotropic intracellular electrical coupling in the 2D and realistic anatomical structure, the 2D model nevertheless represents only a cross-sectional slice through the ventricle; it therefore lacks features of an anatomically realistic 3D ventricular geometry such as the irregular thickness of the wall structure across the entire geometry, layered structure of ventricular tissue and more realistic anisotropy, all of which could influence maintenance of ventricular arrhythmias in the V307L mutation conditions. These potential limitations predicated our additional use of 3D simulations. Though the 3D model incorporates ‘realistic’ ventricular anatomical geometry, it lacks inclusion of a Purkinje fibre network, which may play a role in arrhythmogenesis in the SQT2$^{48}$. In addition, the tissue models in this study do not consider the effect of cardiac mechanics on tissue geometry, which feasibly might influence re-entry$^{41,42}$, particularly for SQTs patients$^{43-45}$.

Whilst it is important that both assumptions and potential limitations of the models used in this study are made explicit, these do not influence fundamentally the conclusions that can be drawn on likely mechanisms by which the KCNQ1 V307L mutation facilitates arrhythmia induction and maintenance, and the possibility of $\text{I}_\alpha$ inhibition as pharmacological modulation in the setting of SQT2. Moreover, it is striking that in our simulations, despite differing levels of complexity, the 2D and 3D simulations yielded qualitatively if not quantitatively similar findings in terms of identifying mechanisms that can account for increased arrhythmia susceptibility in SQT2. This highlights the likely importance of the pro-arrhythmogenic mechanisms identified in this study.

The simulations in this study substantiate that the KCNQ1 V307L mutation is causally linked to QT interval shortening and increased tissue vulnerability to arrhythmogenesis in this form of the SQT2. In addition, we have shown that $\text{I}_\alpha$ blockade normalises the QT interval and successfully terminates re-entry during tachyarrhythmias. Thus, the findings of this study provide a comprehensive explanation for clinical consequences of this form of the SQT2 in terms of abbreviation of repolarisation and susceptibility to arrhythmia and it provides a potential pharmacological approach in designing therapeutic interventions for this form of the SQT2. It is notable that the more recently reported R259H$^{13}$ and F279I$^{46}$ KCNQ1 SQT2 mutations produce similar changes to recombinant channel $\text{I}_\alpha$, to those of V307L KCNQ1. The results reported here may therefore also have relevance to SQT2 associated with these mutations. Additionally, the multi-scale ventricular models developed and employed in this study may have further utility for probing the basis of arrhythmia, both in other forms of the SQT2 and other repolarisation disorders.

Methods

Development of $\text{I}_\alpha$ Markov chain model. The MC model of $\text{I}_\alpha$, incorporating the V307L KCNQ1 mutation was based on the work of Silva and Rudy$^{17}$, who developed it using experimental data at physiological temperature (37°C) including data from human ventricular myocytes representing $\text{I}_\alpha$ activation and deactivation kinetics (MC scheme shown in Supplementary Figure S1). The model was modified to reflect the experimentally observed kinetic properties of WT and V307L mutant “$\text{I}_\alpha$” (KCNQ1 + KCNE1) channels. These kinetic properties include: (i) the profound leftward shift (~36 mV) of channel activation for the V307L mutation compared to WT$^{17}$, (ii) slower channel deactivation in the V307L mutation$^{17}$, (iii) the increased repolarising current during $-36$ mV) of channel activation for the V307L mutation compared to $-40$ mV) of channel deactivation for the V307L mutation compared to $-40$ mV$^{17}$, and (iv) accelerated activation as seen with the V307L mutation conditions. Thus, the key kinetic properties of WT and V307L, we simulated the experimental current-voltage ($\text{I}$-$\text{V}$) relationships for WT and V307L KCNE1 using the voltage clamp protocol from El-Harchi et al.$^{17}$ (Fig. 1Ai–Bi). The membrane potential was held at ~80 mV and then depolarised briefly to ~40 mV for 50 ms, followed by 3 s depolarisations to a range of potentials from ~70 mV to ~60 mV (in 10 mV increments); finally, tail currents were elicited by repolarisation to ~40 mV for 5 s. The currents obtained at the end of the depolarising steps were normalised and compared to experimental data (Fig. 1Ai–Bi). The elicited current traces from the voltage clamp protocol are shown in Fig. 1Ai–Bi.

To obtain the transition rates of the MC model that reproduced the experimentally observed kinetic properties of WT and V307L, we simulated the experimental current-voltage ($\text{I}$-$\text{V}$) relationships for WT and V307L KCNE1 + KCNE1 using the voltage clamp protocol from El-Harchi et al.$^{17}$ (Fig. 1Ai–Bi). The membrane potential was held at ~80 mV and then depolarised briefly to ~40 mV for 50 ms, followed by 3 s depolarisations to a range of potentials from ~70 mV to ~60 mV (in 10 mV increments); finally, tail currents were elicited by repolarisation to ~40 mV for 5 s. The currents obtained at the end of the depolarising steps were normalised and compared to experimental data (Fig. 1Ai–Bi). The elicited current traces from the voltage clamp protocol are shown in Fig. 1Ai–Bi.
By minimising the least-squared difference between the experimental data and the simulation, the variables representing the transition rates that produced the best fit and behaviour of macroscopic currents relative to the experimental data were obtained (Fig. 1Aii–Bii). The minimisation was performed using the Nelder-Mead Simplex algorithm[7]. Relative current proportions for WT and V307L KCNQ1 conditions were then scaled using relative proportions of peak \( I_{KS} \) obtained from AP clamp experiments[7].

**AP clamp and model validation.** The MC model was validated by comparing simulated results with those observed experimentally during ventricular AP clamp (Fig. 1Aiv–Biv). The same digitised ventricular AP used to generate the experimental AP clamp[7] was used in the simulation.

**Heterozygote formulation.** To mimic the heterozygous state of the proband, we constructed a heterozygous formulation (WT-V307L) consisting of 50% WT and 50% V307L channels. We used this formulation to also investigate the effects of the KCNQ1 V307L mutation in this heterozygous condition.

**Cellular and tissue models.** The developed \( I_{KS} \) MC model was incorporated into the 2006 version of the TNNP[9] model of the human ventricular cell AP for consistency with our previous in silico studies on the SQTS[18,19,44]. Single cell models were then incorporated into 1D, 2D and 3D multicellular models with a monodomain tissue representation[18,19,48] of realistic human ventricular geometries. The realistic human geometries were reconstructed as described previously[18,19] with a spatial resolution of 0.2 mm. The anatomical model was segmented into three distinct regions: 25% EPI, 35% MIDDLE and 40% ENDO cells. These proportions are similar to those used in other studies[16,18,21,40] and were chosen as they produced a positive T-wave on the ECG under the WT condition.

Further details regarding multi-scale model development, including methods for simulating the pseudo-ECG, protocols used for measuring restitution of the APD and ERP, initiation of re-entrant excitation waves, dynamics of re-entry, and numerical methods have been documented in detail in our previous studies[16,18,19,44] and are provided in the Supplement.

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Author Contributions
H.Z. and J.C.H. conceived and designed the experiments; A.E.H. and J.C.H. provided in vitro recombinant IKs channel data. I.A. and D.G.W. conducted simulations and prepared figures; I.A., D.G.W., J.C.H., and H.Z. analysed the results. I.A., D.G.W., J.C.H., and H.Z. drafted and edited the manuscript. All authors reviewed the final version of the manuscript.

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