Cellular consequences of HERG mutations in the long QT syndrome: precursors to sudden cardiac death

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Abstract

Background: A variety of mutations in HERG, the major subunit of the rapidly activating component of the cardiac delayed rectifier \(I_{Kr}\), have been found to underlie the congenital Long-QT syndrome, LQT2. LQT2 may give rise to severe arrhythmogenic phenotypes leading to sudden cardiac death. Objective: We attempt to elucidate the mechanisms by which heterogeneous LQT2 genotypes can lead to prolongation of the action potential duration (APD) and consequently the QT interval on the ECG. Methods: We develop Markovian models of wild-type (WT) and mutant \(I_{Kr}\) channels and incorporate these models into a comprehensive model of the cardiac ventricular cell. Results: Using this virtual transgenic cell model, we describe the effects of HERG mutations on the cardiac ventricular action potential (AP) and provide insight into the mechanism by which each defect results in a net loss of repolarizing current and prolongation of APD. Conclusions: This study demonstrates which mutations can prolong APD sufficiently to generate early afterdepolarizations (EADs), which may trigger life-threatening arrhythmias. The severity of the phenotype is shown to depend on the specific kinetic changes and how they affect \(I_{Kr}\) during the time course of the action potential. Clarifying how defects in HERG can lead to impaired cellular electrophysiology can improve our understanding of the link between channel structure and cellular function. © 2001 Elsevier Science B.V. All rights reserved.

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1. Introduction

Increasingly, seemingly idiopathic syndromes are being linked to genetic defects. A recent example is the congenital Long-QT (LQT) syndrome [1–3]. The syndrome is associated with prolongation of the Q-T interval on the ECG, which may precede syncope and sudden cardiac death. In 1995, patients from a family afflicted with a genetically linked form of the Long-QT syndrome (LQT2) were found to have abnormalities in the HERG gene [4]. This gene underlies a potassium channel that is highly expressed in the heart with properties similar to the E-4031 sensitive current [5,6]. Many experiments have since demonstrated that HERG encodes the major subunit of the rapidly activating component of the cardiac delayed rectifier current (\(I_{Kr}\)). Very recently the minK-related peptide 1 (MiRP1) was shown to stably assemble with HERG and give rise to a current strikingly similar to native \(I_{Kr}\) [7].

We develop a structurally based model of cardiac \(I_{Kr}\) in order to elucidate the functional consequences of gene defects in cardiac HERG that lead to abnormalities in repolarization. We have drawn from a variety of experiments presented in the peer-reviewed literature in order to build a robust model that accounts for the breadth of published data. More than 50 mutations in HERG have been linked to the congenital Long-QT syndrome [3]. Defects in the HERG pore have been shown to have heterogeneous cellular phenotypes. Pore mutations may result in a loss of function, sometimes due to a trafficking defect [8], and may or may not coassemble with WT HERG subunits to exert dominant negative effects [9]. Other defects in the channel pore give rise to altered
channel kinetics leading to decreased repolarizing current [10,11]. Nearby mutations in the S4–S5 linker have been shown to variably affect activation [12].

Here, we use Markov models of wild-type (WT) and mutant \( I_{Kr} \) in a cardiac ventricular cell model to evaluate the cellular arrhythmogenic consequences of several HERG mutations associated with LQT2. The Markov models allow for the introduction of state-specific kinetic changes due to mutation induced structural defects. These changes affect adjacent channel states through coupling. We choose as examples two well-characterized defects that alter cell electrophysiology through their effects on channel kinetics. The first, the T474I point mutation, results in an altered voltage dependence of activation coupled with a reduction in macroscopic current density. The second, the R56Q mutation, is a point mutation in the Per–Arnt–Sim (PAS) domain located in the amino-terminus (N-terminus) region of HERG, which normally interacts with the channel and reduces the rate of deactivation [13]. Mutations in this region presumably prevent proper association of the N-terminus with the channel and act to increase the rate of channel deactivation. We choose these two defects as examples of mutations that exert their effects on cellular behavior by reducing the \( I_{Kr} \), repolarizing current through alterations in its channel kinetics.

We have also investigated the effects of the N629D mutation, a ‘gain of function’ defect in HERG that results in loss of C-type inactivation coupled with loss of K\(^+\) selectivity [14]. HERG contains a pore selectivity sequence that is changed by the mutation from GFGN to GFDG, allowing for non-specific passage of monovalent cations. In this study, we incorporate these changes into our model of \( I_{Kr} \) and investigate the effects on the morphology of the action potential and action potential duration.

In addition, we elucidate the effects of a range of reduction in current on the action potential duration (APD), since many HERG defects exert their effects through a dominant negative mechanism, trafficking defects, or reduced mutant protein expression [3]. Finally we evaluate the electrophysiological consequences of a ‘functional knockout’ mutation, where \( I_{Kr} \) current is absent, a situation described clinically in a patient born with such a defect [15]. In all, we provide several examples of heterogeneous genotypes and the mechanism by which each prolongs APD leading to a prolonged QT interval in the ECG, increased dispersion of repolarization, and possibly the development of early afterdepolarizations (EADs) and cardiac arrhythmias.

2. Methods

The general approach to modeling the action potential (AP) is the same as that described for the dynamic Luo–Rudy (LRd) model of a ventricular cell [16] with modifications described in later publications [17,18]. The fast sodium current \( (I_{Na}) \) was recently reformulated using a Markovian approach and incorporated into the whole-cell model [19]. We use a similar approach to reformulate \( I_{Kr} \).

Macroscopic current density is given by:

\[
I_s = G_s \cdot P_{O,s} \cdot (V_m - E_{rev}) \quad \text{where} \quad G_s = \sigma \cdot g_s
\]

\( P_{O,s} \) is the sum of all channel open probabilities, \( V_m \) is the membrane potential, and \( E_{rev} \) is the reversal potential. \( G_s \) is the maximum membrane conductance (channel density (\( \sigma \)) times the unitary channel conductance (\( g_s \))). The channel state probabilities are described by first order differential equations using the methods described in [20]. The model parameters are fit using averaged measured activation, inactivation and deactivation data [5,6,21–23]. Parameters are then optimized to account for the proper behavior of macroscopic currents compared to data recorded at physiological temperatures and ion concentrations from guinea pig ventricular myocytes (see Figs. 3 and 4).

All the simulations were encoded in C/C++. Simulations were implemented (double precision) on a Sun Workstation Ultra 1. A time step of 0.005 ms was used during the stimulus and AP upstroke. At all other times, a 0.01-ms time step was used.

The model for cardiac \( I_{Kr} \) is shown in Fig. 1. The model includes three closed states (C3, C2, C1), an open state (O), and an inactivation state (I). The transition between C2 and C1 is voltage independent [21]. Closed state inactivation (C1 → I) is included from a single closed state, allowing for reduced computation time. The transition rate between C1 and I is the same as from C1 to O [24]. \( I_{Kr} \) exhibits strong inward rectification due to a rapid voltage dependent C-type inactivation at positive membrane potentials [11] (transition from O to I). The transitions between O and I are also dependent on extracellular potassium concentration \( [K^+]_{out} \) [25]. As \( [K^+]_{out} \) is increased the transition rates decrease. In addition, the maximum conductance of \( I_{Kr} (G_{Kr}) \) depends on \( [K^+]_{out} \). These dependences on \( [K^+]_{out} \) are introduced into the model.

Action potential simulations are carried out in epicardial and midmyocardial (M) cells. Epicardial and M cells are simulated by varying the membrane conductance of \( I_{Kr} (G_{Kr}) \) as described previously [18]. Equations of the LRd model can be found in previous publications [16,17,26,27] and can be downloaded from the research section of http://www.cwru.edu/med/CBRTC

3. Results

3.1. Simulated wild type (WT) behavior

The overall result of decreased rates of inactivation and recovery from inactivation at high \( [K^+]_{out} \) is an inactiva-
tion state that is more absorbing. This is demonstrated in the single channel gating of $I_{Kr}$ in Fig. 2 (left panels) and channel mean open times (MOTs) (right panels). Panel A (left) shows experimentally recorded single channel gating of HERG channels at 21°C and 100 mM $[K^+]_{out}$ [24]. Single channels are depolarized from a holding potential of −80 to 100 mV for 300 ms and subsequently hyperpolarized (arrow) to −120 mV for 100 ms (inset, bottom of Fig. 2). Experimental unitary HERG currents (panel A, left) are compared to simulated single channel gating of $I_{Kr}$ (panel B, left) at 100 mM $[K^+]_{out}$ [24]. The simulated single channel gating behavior is at 37°C. The single channel gating behavior in the experiments and the simulations demonstrates channel inactivation from closed states and recovery from inactivation through the open state. Channels may fail to open during depolarization (single channel traces before arrow), due to inactivation from closed states, but conduct an inward current during subsequent hyperpolarization (traces after arrow) due to recovery from channel inactivation through the open state. The experimental traces show a longer latency to first opening compared to the simulations, an expected behavior at the lower temperature under which the experiments are conducted. The model predicts gating behavior at physiological $[K^+]_{out}$ of 4.5 mM in panel C. Here, the inactivation state is less absorbing compared to panels A and B, an overall result of increased rates of inactivation and recovery from inactivation at lower $[K^+]_{out}$. Hence, more channel openings are observed during the depolarization step and the subsequent hyperpolarization (panel C, left). This behavior is consistent with macroscopic current recordings at physiological $[K^+]_{out}$, which have a persistent component of outward current during depolarization (see Fig. 4: experiment, top traces; simulation, bottom traces). This persistent current rapidly reaches a pseudo-steady state due to the rapid equilibrium between open and inactivation states. The MOTs of experimentally recorded HERG channels at $[K^+]_{out}$ of 100 mM and $[K^+]_{out}$ of 4.5 mM (right panels B and C, respectively) are shown for comparison. The simulated MOTs are computed as $1/\Sigma \beta (\text{sum of transition rates leaving the open state, } \beta + \beta')$. At physiological $[K^+]_{out}$ of 4.5 mM, MOTs are shorter due to the increased inactivation rate. This comparison demonstrates that both the simulated and experimentally obtained MOTs exhibit weak voltage dependence over a wide range of voltage regardless of $[K^+]_{out}$ [6,24].

The voltage dependence of activation and rectification (inactivation) of macroscopic $I_{Kr}$ is shown in Fig. 3. The activation curve (panel A) is constructed by normalizing
peak tail current at the indicated test potential to the maximum tail current during repolarization (−100 mV). The solid line is the simulation; filled boxes are experimental data obtained from the E-4031 sensitive current in guinea pig ventricular myocytes [5]. The rectification (R) curve (panel B) is computed as:

\[ R = \frac{\text{peak } I_{Kr}}{\text{peak } I_{Kr\text{, in the absence of rectification}}} \]

Computation of the rectification (inactivation) curve using this method more accurately describes channel inactivation than a standard inactivation protocol since \( I_{Kr} \) availability can be contaminated by deactivation at membrane potentials below 0 mV [7,11]. Rectification is removed in the simulation by making the O→I transition rate equal to zero. The simulated rectification curve is compared to experimental data from Sanguinetti and Jurkiewicz (filled circles) [5]. Rectification is obtained experimentally by extrapolating the linear portion of the peak tail current \( I-V \) curve into the positive potential range [5].

\( G_{Kr} \) is affected by variations in \([K^+]_{out}\). In Fig. 3C, the peak tail current–voltage relationship is shown for \([K^+]_{in}=150 \text{ mM and } [K^+]_{out}=5.4, 50 \text{ and } 150 \text{ mM}.\) Simulated \( I-V \) curves (top) are compared to experimentally recorded rabbit \( I_{Kr} \) (bottom) using the same protocol as in Shibasaki [6]. From a holding potential of −43 mV, the cell is pulsed to the test potential for 500 ms. A subsequent pulse to −43 mV for \([K^+]_{out}=5.4, \text{ and } −73 \text{ mV for } [K^+]_{out}=150 \text{ mM}\).
Fig. 3. Activation (A) and rectification (B) curves are constructed from model $I_{Kr}$ and compared to experimentally recorded guinea pig $I_{Kr}$ at physiological temperature and ion concentrations [5]. The simulated current–voltage ($I$–$V$) relationship at three different $[K^+]_{out}$ is shown (C, top) and compared to experimental data (C, bottom) from Shibasaki [6]. See text for protocol.

$[K^+]_{out}$ = 50 and 150 mM elicits a tail current that is plotted as a function of the test potential. Notice that at $[K^+]_{out}$ = 5.4 mM, the tail current is outward at all membrane potentials ($E_{rev}$ = −89 mV) and exhibits strong inward rectification at positive potentials that compares well with the degree of rectification seen experimentally. At $[K^+]_{out}$ = 50 mM and $[K^+]_{out}$ = 150 mM, the tail current is inward in both the simulation and the experiment due to the equilibrium potentials of −29 and 0 mV at 50 and 150 mM $[K^+]_{out}$, respectively.

Fig. 4 (panel A) contains experimentally recorded $I_{Kr}$ current traces from guinea pig ventricular myocytes [5]. Cells are depolarized for 250 ms from a holding potential of −40 mV and are subsequently repolarized to −40 mV for 750 ms (see protocol, inset). Simulated currents following the same protocol are shown in panel B. The tail current is plotted on a reduced time scale in order to be consistent with the experimental traces [5]. At a test potential of 10 mV or greater, hyperpolarization elicits an outward tail current hook experimentally and in the simulation (arrow). This hook is absent at test potentials 0 mV (arrow) or smaller. The hook reflects recovery from inactivation through the open state. Since recovery from inactivation (I→O) is very fast compared to deactivation (O→C1), a large outward tail current is conducted as channels slowly return to the closed resting states.

3.2. Simulated HERG mutations

The T474I mutation, a point mutation in the S2–S3 linker, alters gating properties and reduces macroscopic $I_{Kr}$ [28,29]. We simulate these changes by altering the voltage dependence of the activation transition rates ($\alpha$ and $\alpha^*$, see Fig. 1 and Appendix) and reducing $G_{Kr}$ by 35%. Fig. 5 compares simulated (panel B) activation and $I$–$V$ curves for WT and T474I mutant $I_{Kr}$ to experimentally measured (panel A) curves. The activation curves (protocol described for Fig. 3) are shown in the left panels. The experimentally measured $V_{1/2}$ for WT and T474I is −15.9 and −43.2 mV, respectively (a shift of −27.3 mV for T474I relative to WT) [28]. The T474I simulated activation curve is shifted by −24 mV ($V_{1/2}$ = −47 mV) relative to the WT curve ($V_{1/2}$ = −23 mV). Experimental recordings were performed at 22–23°C, while the simulations were performed at
physiological temperature (37°C). The leftward shift of the simulated activation curves relative to measured curves is an expected result of the increased temperature [22]. The fully activated $I-V$ relationships of simulated $I_{Kr}$ and experimentally recorded HERG are shown in the right panels of Fig. 5 (protocol described in Zhou et al. [28]). The mutant peak tail current is shifted by $-20$ mV relative to WT in the experimentally recorded and simulated curves. The maximum mutant current is 65% of WT levels.

Mutations in the PAS domain of the amino terminal of HERG act to increase the rate of deactivation [30]. We focus on the R56Q mutation since this mutant can increase the rate of deactivation most profoundly [30]. The mutation is simulated by increasing the $O\rightarrow C1$ transition rate by a factor of 6.3 and the $C2\rightarrow C3$ rate by a factor of 10.5 [30]. These increased rates of deactivation result in a rightward shift in the voltage dependence of activation and rectification (not shown), due to the strong coupling between the discrete states. The simulated kinetic changes result in shifts of $+18$ mV in the activation curve and $+16$ mV in the rectification curve. These shifts are also seen experimentally in HERG channels containing the R56Q mutation expressed in Xenopus oocytes [30] (not shown).

Fig. 6 compares action potentials from WT, T474I and R56Q simulated M-cells. The 1000th AP is shown at a physiological temperature (37°C). The leftward shift of the $G$ (see model in Fig. 1). During the AP upstroke, channels move from the leftmost closed state (C3, panel C) and pass through C2 and C1 (panels D and E, respectively). From the C1 state, channels can pass directly to the open state (O, panel F) or to an unavailable inactivation state I (panel G). Once the channels open, they rapidly inactivate (transition from $O\rightarrow I$). A pseudo-equilibrium is reached between the inactivation transition ($O\rightarrow I$) and recovery from inactivation ($I\rightarrow O$) that begins to favor recovery as the AP plateau repolarizes. As recovery from inactivation becomes more likely, $I_{Kr}$ increases to a pronounced peak (arrow) and contributes to the faster repolarization late in the AP. At this time channels slowly deactivate ($O\rightarrow C1$ transition).

In Fig. 6, the middle panel shows $I_{Kr}$ and associated channel state probabilities during the AP for a T474I mutant M-cell (compare to WT). In addition to the kinetic
Fig. 6. Action potential (AP) of 1000th paced beat of a wild-type (WT) M-cell paced at a cycle length (CL) = 750 ms, is shown in panel A, left. Panel B is the corresponding $I_{K_r}$ during the AP. Panels C–G are the probabilities of residence in the indicated states over the course of the AP (see Fig. 1 for the definition of channel states). The 1000th paced beat of a T474I affected M-cell is shown in the middle panel. Effects of the R56Q mutation on the AP are shown on the right.

Changes, $G_{K_r}$ is reduced by 35% in the simulation. Mutant APD is 16 ms longer than WT. Interestingly, the reduction in $G_{K_r}$, rather than altered channel kinetics, is the major factor in lengthening APD. The alterations in activation kinetics alone have only a minor effect on APD (not shown). The morphology of $I_{K_r}$ early in the AP is altered due to the shift in the voltage dependence of activation. The leftward shift of the activation curve results in early channel openings at less depolarized membrane potentials (panel F, compare to WT, panel F) and an increased rate of $I_{K_r}$ activation (panel B). Channels in the open state are available to inactivate earlier (panel G), resulting in the large notch following activation at the beginning of the current trace (panel B) which reflects fast inactivation. The channel open probability late in the AP (panel F), when $I_{K_r}$ plays a major role in repolarization, displays a pronounced peak (arrow) and is similar to that of WT $I_{K_r}$.

The effect of the R56Q mutation on the cardiac AP is shown in an M-cell in the right panel of Fig. 6. While initial $I_{K_r}$ elicited during the upstroke of the AP is comparable between the WT and R56Q mutant cells (panel B, compare to WT), the late current in the mutant is decreased due to increased probability of the O→C1 transition. This is seen in the open probability during the AP, shown in panel F. In contrast to WT and T474I mutant currents, the R56Q mutant lacks a peak in $P(O)$ late in the AP (panel F, arrow). Deactivation occurs earlier, resulting in less repolarizing current late in the AP where $I_{K_r}$ usually plays a major role in repolarization and determining APD.

Fig. 7 shows the rate dependence of APD in WT (panel A) and the two kinetic mutants T474I (panel B) and R56Q (panel C). There is only a slight prolongation at slow rate in mutant epicardial cells compared to WT (top). However, mutant M cells show significantly greater APD prolonga-
Fig. 7. Rate dependence of APD in WT (A), T474I (B), and R56Q (C) cells. Epicardial cells are shown in the top panels, M-cells in the bottom. M-cells APD is preferentially prolonged at slow rate for both mutations. The table summarizes the rate dependence of APD in M-cells. Values for APD are given in the bottom for each mutant at the different CL.

Fig. 8 demonstrates the effects of the N629D gain of function mutation on the action potential in epicardial (panel A) and M-cells (panels C and E). All cells are paced at a CL = 1000 ms for 1000 beats. The corresponding $I_{Kr}$ is shown beneath each AP trace (panels B, D and F). The mutation is characterized by a loss of C-type inactivation and of ion selectivity, allowing for the passage of Na$^+$ ions in addition to K$^+$ ions. The relative selectivity of the channel for Na$^+$ and K$^+$ ($P_{Na}/P_k$) is 0.65 [14]. We simulate the mutation by eliminating C-type inactivation (O$\rightarrow$I transition rate = 0) and by altering selectivity to the ratio above. Using the Nernst equation, we compute the reversal potential of the N629D $I_{Kr}$ to be $-13$ mV. This reversal potential for mutant $I_{Kr}$ falls within the range of physiological plateau potentials. As a result, early $I_{Kr}$ is an outward repolarizing current carried by K$^+$. However, when the membrane potential repolarizes below $-13$ mV, the current is inward and carried by Na$^+$. This is demonstrated in the $I_{Kr}$ traces during epicardial APs shown in Fig. 8, panel B. During the initial phase of the action potential $I_{Kr}$ is outward, but once the repolarizing membrane potential passes the reversal potential threshold, the current becomes inward (arrows). The inward current late in the plateau results in severe APD prolongation. Even more severe, are the effects of this mutation in M-cells (panel C and E). In panel C, we show an M-cell with $G_{Kr}/G_{Ks} = 7:1$, and in panel E an M-cell with $G_{Kr}/G_{Ks} = 12:1$, both of which fall within the physiological range for currents measured in M-cells. In both M-cells, the inward $I_{Kr}$ carried by Na$^+$ on the background of the smaller $I_{Ks}$ of M-cells, acts to sufficiently prolong the AP plateau to allow for L-type Ca$^{2+}$ channels recovery and reactivation resulting in EADs [26,31]. The morphology of the AP is profoundly disrupted as a result of this mutation.

Fig. 9 demonstrates the effects on the AP of HERG mutations resulting in decreased density of functional channels. We simulate the effect of the resulting current suppression on the morphology and duration of the AP in M-cells. In epicardial cells, these reductions act to prolong APD but do not give rise to the development of EADs (not
Fig. 9. Variable loss of functioning channels can arise from different mutations. Here the effects of a 25, 50 and 75% loss on APD in an M-cell is shown after 1000 paced beats at a CL = 1000 ms. A 25% loss may arise from a reduction in mutant protein expression. The 50 and 75% losses may result from trafficking defects and dominant negative current suppression, respectively. A 75% loss of functional channels results in EADs.

Fig. 10 (panels A and B) demonstrates the effects of changes in pacing rate in a cell with a 50% loss of functional $I_{Kr}$ channels. In epicardial cells (panel A) the resulting reduction in current gives rise to a subtle prolongation of the APD that becomes more pronounced at slow rates. In panel B, the rate-dependent effect in an M-cell is more pronounced. At a slow CL = 2000 ms, the M-cell APD is markedly prolonged and a characteristic ‘shoulder-morphology’ AP is observed.

When $I_{Kr}$ is absent as a consequence of a homozygous mutation (‘functional $I_{Kr}$ knockout’), the patient phenotype is marked by severe prolongation of the QT interval and polymorphic ventricular tachyarrhythmias [15]. In Fig. 10, panels C and D, simulation of this condition results in the most severe disruption of cellular repolarization. Epicardial cells (panel A) have prolonged APD (by 29 ms at CL of 750 ms; 35 ms at a CL of 2000 ms). The 999th and 1000th beats at the indicated cycle length are shown for an M-cell mutation results in removal of C-type inactivation and renders the channel pore non-selective among cations. The inward current (arrows, panel B) is carried by Na⁺ and acts to disrupt the delicate balance of currents during the plateau, shifting it in the inward (depolarizing) direction. Two species of M-cells are shown in panels C ($G_{Kr}$: $G_{Ca} = 7:1$) and E ($G_{Kr}$: $G_{Ca} = 12:1$) with corresponding $I_{Kr}$ shown in D and F, respectively. The mutation profoundly affects the AP morphology in M-cells and leads to EAD formation.

4. Discussion

In this study, functional defects in $I_{Kr}$ due to several HERG mutations are introduced into a comprehensive model of the cardiac ventricular myocyte. The results demonstrate that distinct mutations can have variable effects on current morphology and lead to varying degrees of altered electrophysiology, depending on kinetic changes induced by the mutation. The use of a Markovian model to represent $I_{Kr}$ deviates from the traditional Hodgkin–Huxely approach [34]. The Markovian scheme represents distinct channel states and coupling between these states, allowing us to relate state-specific kinetic properties of ion channels to the electrophysiological behavior of the whole cell.

Heterogeneous mutations in HERG that give rise to the same syndrome, congenital LQTS, may differ in severity
and differentially affect the morphology of $I_{Kr}$ during the AP. The T474I mutation predominantly affects the current at the onset of the AP through a leftward shift in the voltage dependence of activation. However, recovery from inactivation and deactivation are unaffected. Hence, the morphology of the late current is preserved in the T474I mutation, as evidenced by the preservation of the pronounced peak in channel open probability (P(O)) late in cellular repolarization seen in WT cells (Fig. 6, arrow). The result is a relatively small effect on APD.

In contrast, the R56Q mutation affects deactivation kinetics that decrease $I_{Kr}$ late in the AP. The delicate equilibrium between recovery from inactivation and deactivation is altered as a result of this defect. The increased rate of deactivation acts to ‘pull’ channels from the open state into closed states before they can conduct a substantial current. It is clear from examination of the current during the AP (Fig. 6, right), that late $I_{Kr}$ is diminished due to the lack of the late maximum in channel P(O), which markedly slows AP repolarization and prolongs APD. Here, we focus on the most severe of the PAS mutations, but this behavior can be generalized to various defects that act to speed deactivation [30].

The N629D mutation gives rise to $I_{Kr}$ that fails to inactivate. This gain of function would seemingly manifest as a larger $I_{Kr}$ that would act to shorten APD. However, the disruption of the pore selectivity sequence that also results from the mutation allows for passage of Na$^+$ ions into the cell. The loss of ion selectivity gives rise to a shift in the reversal potential of $I_{Kr}$ to $-13 \text{ mV}$. This elevated reversal potential is crossed during the plateau repolarization phase of the action potential. Once the membrane potential falls below $-13 \text{ mV}$, the channel passes an inward current carried by Na$^+$ (see Fig. 8, panel B, arrows). This inward current acts to depolarize the membrane late in the AP and prolongs APD. It results in a severe cellular phenotype similar to the effects of the $\Delta$KPQ gain of function mutation described previously in the cardiac Na$^+$ channel [19]. We also investigated the effects of loss of C-type inactivation and of loss of ion selectivity individually (not shown). Loss of inactivation caused a reduction in APD due to increased $I_{Kr}$, while loss of ion selectivity was sufficient to cause APD prolongation.

To further explore the effects of HERG mutations, we have characterized the effects of a range of reductions in $G_{Kr}$ on the morphology of the AP. A 25, 50, or 75% reduction in current could arise in a heterozygous mutation from reduced protein expression, processing or transport defect, or from dominant negative effects, respectively [15,28,32]. A 50% reduction in $I_{Kr}$ acts to prolong the epicardial AP slightly, but has a major prolongation effect.
on the APD of M-cells especially at slow rates. A functional knockout (homozygous mutation) results in the most severe disruption of cellular repolarization, consistent with patient phenotypes observed clinically. The mutation acts to prolong APD at plateau potentials, providing sufficient time for reactivation of the L-type Ca$^{2+}$ current, which depolarizes the cell and generates the EAD [26,31]. It should be emphasized that L-type reactivation during the prolonged plateau is common to all mutations investigated in this paper and provides the depolarizing charge during the EAD upstroke.

In general, structural and functional information is obtained from isolated membrane patches and/or cloned channels in expression systems (e.g. *Xenopus* oocytes) removed from the environment where they function physiologically. The natural cellular milieu is highly interactive and modulates single channel kinetic behavior. Cellular models are unique tools that allow for reinstitution of single channels into the cell in order to investigate integrated electrophysiological behavior. That the physiological effect of a given channel defect depends on the cellular environment is demonstrated here by the greater severity of APD prolongation and disruption of repolarization in M cells, suggesting that such mutations can increase APD differences and dispersion of repolarization in the myocardium [33]. A substrate with increased dispersion of repolarization is susceptible to the development of unidirectional conduction block and reentrant arrhythmias [2,35]. Such arrhythmias can be triggered by the EADs associated with the mutations which provide premature stimuli to a highly susceptible substrate [36].

In the context of arrhythmia, it is interesting to note that arrhythmias associated with HERG mutations and LQTS are not rate dependent to the same extent as other forms of inherited LQTS [37]. Our simulations predict that HERG defects can lead to the development of cellular arrhythmogenic responses (APD prolongation and early afterdepolarizations) at slow rates in the absence of neural influences and may explain the development of arrhythmias in patients during sleep and relaxation. However, HERG defects have also been linked to arrhythmias in patients startled by loud noise, such as a telephone or alarm clock, suggesting a role for the autonomic nervous system in increasing susceptibility to arrhythmic events [38,39]. Previous modeling studies in our laboratory have incorporated the effects of β-adrenergic stimulation in a model of the WT cell [31]. Future efforts will expand the previous studies to investigate effects of the autonomic nervous system in the context of channelopathies including LQTS.

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### Appendix. I$_{Kr}$ formulation

$$I_{Kr} = G_{Kr} \cdot P(O) \cdot (V_{in} - E_K)$$

$P(O) = $ open probability of $I_{Kr}$

$$G_{Kr} = 0.0135 \cdot [K^+]_{out}^{0.59}$$

$$E_K = (R \cdot T/F) \cdot \ln([K^+]_{out}/[K^+]_{in})$$

### Wild-type rate constants

- $C_1 \rightarrow O$ or $C_1 \rightarrow I \quad \alpha_\alpha = 65.5 \cdot 10^{-3} \cdot e^{(0.05547153 \cdot (v - 36))}$
- $C_2 \rightarrow C_1 \quad \alpha_{in} = 2.172$
- $C_3 \rightarrow C_2 \quad \alpha = 55.5 \cdot 10^{-3} \cdot e^{(0.05547153 \cdot (v - 12))}$
- $C_2 \rightarrow C_3 \quad \beta = 2.357 \cdot 10^{-3} \cdot e^{(-0.036588 \cdot (v))}$
- $C_1 \rightarrow C_2 \quad \beta_{in} = 1.077$
- $O \rightarrow C_1 \quad \beta \beta = 2.9357 \cdot 10^{-3} \cdot e^{(-0.02158 \cdot (v))}$
- $I \rightarrow O \quad \alpha_i = 0.439 \cdot e^{(-0.02352 \cdot (v + 25))} \cdot 4.5/[K^+]_{out}$
- $O \rightarrow I \quad \beta_i = 0.656 \cdot e^{(0.000942 \cdot (v))} \cdot 4.5^{0.3}/[K^+]_{out}^{0.3}$
- $I \rightarrow C_1 \quad \mu = (\alpha_i \cdot \beta \alpha \cdot \alpha \alpha)/(\alpha \alpha \cdot \beta \beta)$

### T474I rates

- $C_1 \rightarrow O \quad \alpha_\alpha = 65.5 \cdot 10^{-3} \cdot e^{(0.05547153 \cdot (v + 25))}$
- $C_3 \rightarrow C_2 \quad \alpha = 55.5 \cdot 10^{-3} \cdot e^{(0.05547153 \cdot (v + 6))}$

### R56Q rates

- $C_2 \rightarrow C_3 \quad \beta = 2.357 \cdot 10^{-3} \cdot e^{(-0.036588 \cdot (v))} \cdot 10.5$
- $O \rightarrow C_1 \quad \beta \beta = 2.9357 \cdot 10^{-3} \cdot e^{(-0.02158 \cdot (v))} \cdot 6.3$

### N629D rates

- $O \rightarrow I \quad \beta_i = 0.0$
- N629D loss of selectivity
Na:K permeability \( (P_{Na}/P_{K}) = 0.65 \)

\[
E_K = (R \cdot T/F) \\
\quad \cdot \ln([K^+]_{out} + P_{Na}/P_{K} \cdot [Na^+]_{out}/[K^+]_{in} + P_{Na}/P_{K} \cdot [Na^+]_{in})
\]

References


