Subunit Interaction Determines $I_{Ks}$ Participation in Cardiac Repolarization and Repolarization Reserve

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**Background**—The role of $I_{Ks}$, the slow delayed rectifier K$^+$ current, in cardiac ventricular repolarization has been a subject of debate.

**Methods and Results**—We develop a detailed Markov model of $I_{Ks}$ and its $\alpha$-subunit KCNQ1 and examine their kinetic properties during the cardiac ventricular action potential at different rates. We observe that interaction between KCNQ1 and KCNE1 (the $\beta$-subunit) confers kinetic properties on $I_{Ks}$ that make it suitable for participation in action potential repolarization and its adaptation to rate changes; in particular, the channel develops an available reserve of closed states near the open state that can open rapidly on demand.

**Conclusions**—Because of its ability to form an available reserve, $I_{Ks}$ can function as a repolarization reserve when $I_{Ks}$, the rapid delayed rectifier, is reduced by disease or drug and can prevent excessive action potential prolongation and development of arrhythmogenic early afterdepolarizations. (Circulation. 2005;112:1384-1391.)

**Key Words:** action potentials ■ electrophysiology ■ ion channels

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Mutations to the cardiac potassium channel gene KCNQ1 (KvLQT1) have been linked to the long QT syndrome LQT1, which predisposes patients to arrhythmia during exercise and emotional stress, conditions that involve high levels of $\beta$-adrenergic stimulation. KCNQ1 is a 6-transmembrane domain protein that can form functional homomeric potassium channels and can also coassemble with the single transmembrane protein KCNE1 (MinK).1 Together these gene products reconstitute the $I_{Ks}$ channel. Mutations to KCNE1 have also been linked to LQT (LQT5).2 In addition, transmural heterogeneity of $I_{Ks}$ expression in ventricular myocardium gives rise to mid-myocardial cells (M cells) with a longer action potential (AP) duration (APD) and greater APD rate adaptation than epicardial or endocardial cells in many species.3 $I_{Ks}$ is augmented by $\beta$-adrenergic stimulation,4 suggesting its important role in mediating cardiac electrophysiological response.

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These findings suggest that $I_{Ks}$ is important for AP repolarization and APD adaptation to changes in rate, as demonstrated in guinea pig.5–7 However, $I_{Ks}$ density has been reported to be much lower in larger mammals, specifically in canine and human ventricle.8 In canine myocytes, L-type calcium current, an inward depolarizing current, has been shown to mediate APD rate adaptation under control conditions without $\beta$-adrenergic effects.7 However, AP repolarization requires a sufficient outward repolarizing current during phases 2 and 3 of the AP. Such current is carried by $I_{Ks}$ (the rapid delayed rectifier) and $I_{Kr}$, with $I_{Ks}$ playing a primary role in large mammals under normal physiological conditions and in the absence of $\beta$-adrenergic stimulation. Because phases 2 and 3 depend on a delicate balance between inward and outward currents, one cannot rule out a priori an important role for $I_{Ks}$. Moreover, the arrhythmic consequences of LQT1 and LQT5 mutations, the existence of only 2 repolarizing currents ($I_{Ks}$ and $I_{Kr}$) that constitute the delayed rectifier, and incorporation of the $\beta$-adrenergic signaling molecules into the $I_{Ks}$ channel complex strongly suggest an important role for $I_{Ks}$ in human heart electrophysiology under various conditions. It has been hypothesized that in large mammals $I_{Ks}$ constitutes a “repolarization reserve” (RR) that compensates for reductions in other repolarizing currents, in particular $I_{Ks}$, caused by mutations (hereditary LQT2) or drugs (acquired LQT syndrome).9,10 There is growing consensus that in the absence of RR, certain drugs such as sotalol (antiarrhythmic), erythromycin (anti-infective), chlorpromazine (antipsychotic), and methadone can trigger a life-threatening arrhythmia.9 The possibility of $I_{Ks}$ generating this reserve and the dependence of $I_{Ks}$ participation on its kinetics remain to be elucidated.

In the present study we examine the hypothesis that $I_{Ks}$ can participate in AP repolarization because of kinetic properties conferred by interaction between its KCNQ1 and KCNE1 subunits. We present detailed, experimentally based Markov...
models of KCNQ1 and \( I_{K_s} \) and examine their kinetic behavior during the AP at slow and fast rates. By comparing KCNQ1 behavior with \( I_{K_s} \), we isolate the effect of the modulatory KCNE1 subunit on the AP. Results show that because of its kinetic properties, \( I_{K_s} \) can create an available reserve (AR) of channels at fast rates that can open and generate a larger repolarizing current. In the presence of \( I_{K_s} \) block, this AR prevents excessive APD prolongation and the formation of arrhythmogenic early afterdepolarizations (EADs). Such properties are not present in homomeric KCNQ1 channels and therefore require interaction with KCNE1. The AR concept relates to reserve within a single channel, thereby extending the RR concept that involves compensation for one repolarizing current by another.

**Methods**

Markov models of KCNQ1 and \( I_{K_s} \) are derived from experimental data and published \( K^+ \) channel models. Koren et al.\(^{11}\) described the \( K^+ \) delayed rectifier RCK1 with a Markov model of 4 independent voltage sensor transitions and 1 cooperative voltage-independent transition before the open state. Zagotta et al.\(^{11}\) expanded this model to study Shaker \( K^+ \) channels by assuming that each voltage sensor undergoes 2 conformational changes before channel opening and successfully reproduced delayed activation (sigmoidal activation). A delay of several milliseconds has also been observed for KCNQ1\(^{11}\) and \( I_{K_s} \) activation, suggesting that at least 2 voltage sensor transitions occur before channel opening. Experiments (J. Cui, PhD, personal communication, November 2004) also suggest a voltage-independent transition before the open state, as proposed by Koren et al.\(^{11}\) for RCK1.

The Markov schemes we developed for KCNQ1 and \( I_{K_s} \) are shown in Figure 1A and 1B. Two closed-state zones are shown; green represents states in which at least 1 voltage sensor must still make a first transition, and blue represents states in which only faster second transitions are necessary for opening. Model derivation is described in the online-only Data Supplement. Criteria for fitting parameters are described below.

The KCNQ1 model is based on frog oocyte recordings\(^{11,12}\) (Figure 2). Simulated activation to various potentials (Figure 2A), inactivation measured by a triple-pulse protocol (Figure 2C), and dependence of deactivation time constant on prepulse duration (Figure 2E) were included in the fitting procedure (details are in the online-only Data Supplement). The reversal potential was \(-50 \text{ mV} \) on the basis of the current-voltage relationship that shows no activation at \(-10 \text{ mV} \) (Figure 2B). Therefore, instantaneous current (possibly due to leakage) was subtracted. Current tracings were normalized according to the current-voltage relationship curve so that the steady state value for each tracing would match the experimental average. Deactivation at \(-50 \text{ mV} \) was fit directly to published data.\(^{15}\) Rate of deactivation at resting \( V_m \) and current accumulation at fast pacing rates were simulated with \( I_{K_s} \) inserted in the whole-cell model (see below); comparison with experimental data\(^{15}\) is shown in Figure 3B. Additionally, mean current increase at fast rates and dynamic conductance (corresponding to APs and \( I_{K_s} \) channels) were used (ratio of \( Na^+ : K^+ \) permeability of 0.01833).\(^6\)

Human \( I_{K_s} \) activation kinetics (Figure 3C) were fit to data from Kupershmidt et al.,\(^{17}\) who recorded expressed human \( I_{K_s} \) in HEK cells at \( 37^\circ \text{C} \). [Na\(^+\)]\(_o\) was 145 mmol/L and [K\(^+\)]\(_o\) was 4 mmol/L. Expressed currents were fit because they display activation kinetics more clearly than currents recorded from native cells while still displaying activation at \(-10 \text{ mV} \) and a delay before slow activation, as in native cells.\(^8\) Deactivation was constrained by time constants measured in human ventricular myocytes at \( 37^\circ \text{C} \) for [Na\(^+\)]\(_o\)=144 mmol/L and [K\(^+\)]\(_o\)=4 mmol/L to ensure that open-state accumulation was not overestimated.\(^8\) Reversal potential was calculated as for guinea pig \( I_{K_s} \). Comparisons with experimental data are shown in Figure 3D.

Markov models for \( I_{K_s} \), \( I_{K_s} \) (updated version), and \( I_{K_s} \) were inserted in the Luo-Rudy (LRd) model of the guinea pig ventricular cell. AP simulation conditions are discussed in the online-only Data Supplement.

**Results**

**KCNQ1 Model Validation and Current Properties**

The KCNQ1 model (Figure 1A) was optimized to reproduce experimental protocols designed to isolate the essential channel features.\(^{13}\) Sigmoidal activation is reproduced with multiple closed-state transitions before opening. This allows for minimal initial current followed by a steep rise after \( \approx 5 \text{ ms} \), as observed experimentally\(^{13}\) (Figure 2A). Two time constants of activation (fast followed by slow) are evident, with a slow rise in current observed even at 2-second depolarization, the last time point for which the current-voltage relationship is experimentally measured (Figure 2A, 2B).
Next, the triple-pulse protocol (Figure 2C, 2D), which measures channel inactivation at different potentials, is simulated. A short hyperpolarizing pulse to \(-130\) mV allows most channels to recover from inactivation while preventing significant deactivation (Figure 2C). In the model, 5 open states allow channels to occupy open states far from the closed state (O4 and O5, Figure 2C), which delays deactivation during the hyperpolarizing pulse. However, transitions toward the closed state are rapid enough to reproduce the measured time constant of deactivation at \(-70\) mV (Figure 2D; simulated \(\tau_{\text{deact}}\) is 377 ms, measured \(\tau_{\text{deact}}\) is 348 ms). Peak current and time constant of inactivation during the following (third) pulse are consistent with experimental results13 (Figure 2D), demonstrating rapid and voltage-dependent inactivation.

Finally, the dependence of deactivation time constant and relative inactivation (defined in the online-only Data Supplement) on pulse duration are simulated. KCNQ1 deactivation rate varies with pulse duration. As channels enter open states that are farther from the closed states, the rate of entry back into the closed states (\(\tau_{\text{deact}}\)) is slowed (Figure 2F). \(\tau_{\text{deact}}\) is greater than estimated experimentally from the dependence of deactivation on pulse duration (Figure 2E inset). However, a different experimental protocol (voltage dependence of deactivation) in the same study13 estimates \(\tau_{\text{deact}}\) close to the model-determined value at \(-60\) mV (502±27 ms for experiment; 594 ms for model). The different \(\tau_{\text{deact}}\) from the 2 experimental protocols may be explained by the need to compensate for the inactivation hook in the pulse-duration protocol (Figure 2E inset) when fitting an exponential, a procedure that can have a major effect on the estimate of \(\tau_{\text{deact}}\). In the model such correction is not necessary because deactivation is measured directly as the decay of occupancy in the open and inactivated states.

In contrast to rapid inactivation during the third pulse (above), a single depolarizing pulse from \(-80\) mV to 20 mV results in slow onset of inactivation (>200 ms pulse duration needed to elicit hook in tail current; Figure 2E) because of multiple transitions through open states. These transitions also govern relative inactivation (protocol not included in optimization), which shows an experimentally observed 75-ms delay13 (Figure 2F).

The \(I_{Ks}\) model resembles the KCNQ1 model closely, with the number of open-state transitions truncated and inactivation...
removed (Figure 1). Significant differences are introduced by changes in the transition rates. For example, increased KCNE1 mRNA in frog oocytes results in greater delay before activation. This subunit effect is simulated by decreasing the transition rate from the first (resting) voltage sensor position in the heteromeric guinea pig $I_{Ks}$ channel to half the homomeric channel rate, confining more channels to zone 2 (Figure 3A, 3C insets). Once channels transition out of this zone, they activate rapidly. However, some channels remain in zone 2, causing a slow rise in current that continues even after several seconds (Figure 3A).

The first voltage sensor position is more stable for human $I_{Ks}$ than KCNQ1 (C1 occupancy is 9 times greater at 80 mV), resulting in slower activation and a continuous current increase even after a 5-second pulse to 60 mV. Once channels transition into O1, they rapidly enter O2 (at 40 mV, the rate into O2 is 6.6 times faster than C1 to O1). The presence of 2 open states allows for simultaneous reproduction of fast activation, steady state current-voltage relationship, and slow deactivation by facilitating slow deactivation without requiring slow activation to reproduce the steady state current.

$I_{Ks}$ Role in Rate Adaptation of APD

Simulated whole-cell APs computed with the guinea pig $I_{Ks}$ Markov model are shown at fast and slow pacing rates (Figure 4A). At the fast rate, peak $I_{Ks}$ increases moderately compared with the slow rate (Figure 4B); however, the more rapid rise of $I_{Ks}$ at the fast rate leads to greater repolarizing

**Figure 3.** Guinea pig and human $I_{Ks}$. A, Simulated guinea pig $I_{Ks}$ current resulting from protocol in inset. A delay of several milliseconds is observed before depolarization at voltages <20 mV and is inversely proportional to voltage (inset enlarges the initial portion of the current tracings). Tail currents show no hook, indicating absence of inactivation, as in experiment. B, Simulated steady state guinea pig current-voltage (I-V) relationship from the protocol in A is compared with experiment (circles). Adapted from reference 15 with permission from Lip-pincott Williams and Wilkins. C, Human $I_{Ks}$ current resulting from the protocol in inset. Activation is slower than guinea pig $I_{Ks}$ with a longer (20 ms) delay, and tail currents deactivate faster, as observed experimentally. D, Simulated steady state I-V relationship for human $I_{Ks}$ compared with experiment (circles, protocol in C) and time constant of deactivation time constant compared with experiment (squares, protocol in inset). Adapted from references 8 and 17, copyright 2001 and 2002, with permission from The European Society of Cardiology.

**Figure 4.** $I_{Ks}$ role in APD adaptation. A, Fortieth AP at CL=250 ms (thin line) and CL=1000 ms (thick line) under control conditions. B, Guinea pig $I_{Ks}$ during the APs in A. C, State occupancy during the 40th AP at slow rate (1000 ms). Green shows occupancy in zone 2 (Figure 1). Blue shows occupancy in zone 1. Red shows open channel occupancy (O1+O2). At AP initiation, most channels reside in zone 1. Accumulation in zone 1 facilitates channel opening and larger $I_{Ks}$ during the AP (B), causing APD shortening (A). Buildup of zone 1 reserve underlies the participation of $I_{Ks}$ in APD rate adaptation. Open-state accumulation is minimal.
cause APD shortening at fast rates (adaptation). Channels accumulate in zone 1 between APs, allowing 1 before opening. In contrast, at fast rates (Figure 4D) and the open states O1 and O2 (red) (Figure 1). During slow transitions rather than its open-state accumulation. Figure 4C shows total occupancy in zone 2 (green), zone 1 (blue), transients that measured a 28% increase at these rates.16 This is a conservative value compared with AP clamp at cycle length (CL)

The Ik1 mediation of adaptation is a result of its closed-state transitions rather than its open-state accumulation. Figure 4C and 4D show total occupancy in zone 2 (green), zone 1 (blue), and the open states O1 and O2 (red) (Figure 1). During slow pacing (Figure 4C), most Ik1 channels reside in zone 2 before AP depolarization and must make a slow transition into zone 1 before opening. In contrast, at fast rates (Figure 4D) channels accumulate in zone 1 between APs, allowing Ik1 to activate rapidly via the second transition. There is not sufficient time between beats for the channels to transition back to zone 2 before the next AP, as at slow rates. Although there is some open-state accumulation due to slow deactivation, its effect on the current is opposed by the lower driving force resulting from a lower peak membrane potential at fast rates. Consequently, the primary mechanism for adaptation is the accumulation of closed states in zone 1, where a fast transition to the open states generates a rapid Ik1 rise during phase 1. Thus, at fast rates an AR is built ready to open very rapidly on demand. The overall low probability of channels residing in open states at slow and fast rates (Figure 4C, 4D) indicates that many channels are closed during the AP, creating a large Ik1 reserve.

AP clamp experiments have measured Ik1 conductance (gKs) during the AP at fast and slow rates, showing that open-state accumulation is minimal (Figure 5, bottom panel, arrows). Rather, Ik1 activates more rapidly at a fast rate than at a slow rate, as also observed in our simulations (Figure 5, top panel). The faster activation at fast pacing in experiments can be explained by the accumulation in closed states near to the open state that are readily available to open at the fast rate, as simulated by the model (Figure 4). Thus, the experiments support the concept of AR as a mechanism for Ik1 participation in repolarization and its dependence on rate.

Human Ik1 activates more slowly than guinea pig Ik1, and deactivates more rapidly (Figure 3C; compare with Figure 3A). The guinea pig cell provides a natural model for testing Ik1 in an environment in which Ik1 plays only a secondary role in repolarization. To evaluate the capability of human Ik1 to serve as a RR and to compensate for reduced Ik1, we inserted a model of human Ik1 in the guinea pig LRd model (Figure 6). As can be seen in Figure 6F (solid line), with increased maximum conductance, the Ik1 in the guinea pig cell can generate APD adaptation (gKs ~4 times greater than guinea pig to obtain 200 ms APD). In contrast, when homomeric KCNQ1 channels are inserted, there is less APD adaptation at fast rates (Figure 6F, dashed line). This attenuation of APD adaptation is surprising because the deactivation kinetics of KCNQ1 are slower than human Ik1, allowing for open-state accumulation (Figure 6D). However, Ik1 channels accumulate in zone 1 (Figure 6E, black), which is composed of closed states that only require rapid transitions to open, allowing for rapid activation that leads to large current during the repolarization phase of the AP (Figure 6C, thin line). This large Ik1 current during repolarization proves more important for shortening APD than the instantaneous current generated by KCNQ1. As shown earlier (Figure 4), zone 1 accumulation rather than open-state accumulation underlies the increase of Ik1 at fast rates. Such AR closed-state accumulation in zone 1 does not occur in KCNQ1 channels (Figure 6E, gray). Instead, a large percentage of channels activate even at slow rates (open-state occupancy at CL = 10,000 ms is 30% for KCNQ1 versus 7% for Ik1), preventing an increase at fast rates.

Ik1 and KCNQ1 During Ik1 Block

Figure 7A shows mean Ik1 (black) compared with KCNQ1 current (gray) during the first and 40th APs at CL = 500 ms. Ik1 increases ~50% from the first to the 40th AP, whereas KCNQ1 remains almost the same over 40 paced beats (inset shows current tracings for the first 9 beats). When Ik1 is blocked, greater Ik1 increase is observed over 40 paced beats (relative to control conditions), whereas only a limited increase is noted for KCNQ1. The difference between Ik1 and KCNQ1 behavior in the presence of Ik1 block is due to the nature of current accumulation. Ik1 block prolongs the AP plateau and elevates its potential, thus enhancing transitions toward the open states. KCNQ1 accumulation is mostly in the

**Figure 5.** Dynamic guinea pig Ik1 current during AP clamp. Bottom, Experimentally measured Ik1 conductance (Chromanol 293B-sensitive current, gKs) at fast (CL = 250 ms) and slow rates (CL = 1000 ms) (thick and thin lines are mean and confidence limits, respectively). Arrows indicate open-state occupancy at AP onset. Reprinted from reference 16 with permission from Blackwell Publishing. Top, Simulation also shows little open-state accumulation (arrow) even at fast rates; instead, in both simulation and experiment an increase in Ik1 activation rate during fast pacing is responsible for greater Ik1 during the AP.
open state, which is at the expense of populating closed states that constitute the AR. Consequently, most channels open on every beat, allowing for only limited accumulation between beats. In contrast, \( I_{Ks} \) shows minimal open-state accumulation, and \( I_{Ks} \) block facilitates transitions into closed states that constitute the AR, allowing for greater \( I_{Ks} \) accumulation over many beats.

When a pause is simulated after 40 beats in the presence of \( I_{Kr} \) block (Figure 7B), the postpause AP with KCNQ1 develops an EAD as a consequence of insufficient KCNQ1 current late during the AP. In contrast, the AP with \( I_{Ks} \) repolarizes normally because of the superior ability of \( I_{Ks} \) to activate during phase 3 of a prolonged AP.

**Discussion**

We show that interaction of KCNQ1 with the \( \beta \)-subunit KCNE1 to form \( I_{Ks} \) alters kinetics so that an AR is created at fast pacing rates. In contrast to KCNQ1, which does not generate a significant AR, \( I_{Ks} \) causes greater APD adaptation and protects against EADs when \( I_{Ks} \) is reduced.

The AR concept can be examined experimentally. For example, a gapped double-pulse protocol with variation in the gap width could be used to characterize the AR in both guinea pig and human ventricular myocytes. In this protocol, a depolarizing pulse to 40 mV for 2 seconds would activate the channel. This step would then be followed by a repolarizing step of variable duration (from 10 ms to 1 second) to 80 mV so that the channel could partially deactivate. A second depolarizing pulse to 40 mV would be used to measure the rate of activation, providing a measure of the AR.

Our simulations also predict that \( I_{Ks} \) will increase more at fast rates when \( I_{Kr} \) is blocked (Figure 7), a manifestation of AR. This prediction could be experimentally tested by blocking \( I_{Kr} \) and measuring the increase in \( I_{Ks} \) during an AP clamp at slow and fast rates. Another protocol to test interaction between \( I_{Ks} \) and \( I_{Kr} \) would be to reduce \( I_{Ks} \) and measure the change in APD from control during pacing, and then to apply \( I_{Kr} \) reduction to control and block \( I_{Ks} \) to the same degree as in the previous protocol. Our simulations predict that the same \( I_{Ks} \) reduction will have a much greater effect on APD when \( I_{Kr} \) is also reduced.

At fast rates, the ability of guinea pig \( I_{Ks} \) to accumulate between APs allows it to participate in APD shortening. However, in human myocytes \( I_{Ks} \) deactivates rapidly, bring-
significant AR. Consequently, the KCNQ1 AP does not adapt the open states because of slow deactivation but show no behavior of homomeric KCNQ1 channels that accumulate in $I_{\text{Kr}}$ for time (dependence over the first 9 APs is shown in inset). When $I_{\text{Kr}}$ is blocked (right), $I_{\text{Kr}}$ increases further (doubling compared with first control AP), and KCNQ1 increase is small. B, In presence of $I_{\text{Kr}}$ block, pause after 40 APs at CL=500 ms is simulated with $I_{\text{Kr}}$ or KCNQ1. Postpause AP with $I_{\text{Kr}}$ shows normal repolarization; AP with KCNQ1 develops a pause-induced EAD.

This novel mechanism of adaptation contrasts with the behavior of homomeric KCNQ1 channels that accumulate in the open states because of slow deactivation but show no significant AR. Consequently, the KCNQ1 AP does not adapt as much as the $I_{\text{Kr}}$ AP (Figure 6). The fundamental difference between $I_{\text{Kr}}$ and KCNQ1 channels is the degree of participation of the first voltage sensor transition in channel activation. Because of the stabilization of the voltage sensor first position by KCNE1, only a small fraction of $I_{\text{Kr}}$ channels activate at slow rates, creating a reserve of channels that can activate at fast rates to accelerate repolarization. In contrast, a large percentage of KCNQ1 channels activate even at slow rates, preventing buildup of such a reserve. As shown in Figure 6C, the initial jump (arrow) of $I_{\text{Kr}}$ current that reflects open-state accumulation at fast rates is small and is followed by a steady increase to a peak of 3.4 $\mu$A/μF late during the AP. At this phase of the AP, $I_{\text{Kr}}$ has a maximal effect on repolarization and APD. In contrast (Figure 6D), KCNQ1 current displays a large initial jump (arrow) due to large open-state accumulation but does not increase during the AP in the absence of an AR; its magnitude stays at $\approx 2 \mu$A/μF and lacks the late peak that is important for repolarization. Thus, interaction with the KCNE1 subunit strongly influences the $I_{\text{Kr}}$ profile during the AP; it augments AP shortening and is essential to normal $I_{\text{Kr}}$ function and its participation in APD adaptation. Other differences between KCNQ1 and $I_{\text{Kr}}$, such as open-state behavior and flickery block, are discussed in the online-only Data Supplement.

As stated earlier in this report, $I_{\text{Kr}}$ is not likely to participate in rate adaptation in large mammals under control conditions. However, when $I_{\text{Kr}}$ is reduced, $I_{\text{Kr}}$ is the only remaining major repolarizing current. The guinea pig myocyte provides a natural electrophysiological environment in which $I_{\text{Kr}}$ is reduced. Under these conditions, we show that human $I_{\text{Kr}}$ can mediate rate adaptation when the maximum conductance of the current is increased (Figure 6). $\beta$-Agonists, which are present in the normal physiological environment even at basal heart rates, can readily confer such an increase in $I_{\text{Kr}}$ conductance.

In the case of pathologically reduced $I_{\text{Kr}}$ (by a mutation or a drug), outward currents carried by other channels (RR) can prevent excessive AP prolongation, EADs, and triggered activity. It is hypothesized that $I_{\text{Kr}}$ reduction in conjunction with a compromised RR is a precursor to arrhythmia, especially after a pause. To test the ability of $I_{\text{Kr}}$ to participate in the RR, we compared $I_{\text{Kr}}$ with KCNQ1 accumulation under control conditions and with $I_{\text{Kr}}$ block. During pacing, $I_{\text{Kr}}$ displays greater accumulation than KCNQ1 and increases further when $I_{\text{Kr}}$ is reduced (Figure 7A). When the pause protocol is simulated with blocked $I_{\text{Kr}}$, increased $I_{\text{Kr}}$ results in normal repolarization, whereas the postpause AP with KCNQ1 develops an EAD (Figure 7B). These simulations indicate that the ability of $I_{\text{Kr}}$ to form an AR, which leads to accumulation at fast rates, makes it an ideal candidate to play a critical role in RR. The accepted RR concept involves compensation for one repolarizing current by another. This study demonstrates the existence of a reserve (AR) within a single channel ($I_{\text{Kr}}$), thereby extending the RR concept to include single-current reserve.

We describe $I_{\text{Kr}}$ with 2 transitions, a slow transition (to zone 1) followed by a fast transition (to open), which implies...
that a 2-closed-state model (rather than 15 states) representing lumped voltage sensor transitions could adequately describe $I_{\rm Ks}$ activation. However, this model would not reproduce the experimentally observed delay before activation,\textsuperscript{14,19} which has important consequences during the AP. To reproduce these kinetics, a semi-Markov 2-closed-state model could be used that would introduce a memory property to the channel via time-dependent transition rates. Such models have been proposed\textsuperscript{20} but introduce another level of complexity with the addition of memory. The present model was chosen because of its correlation to the tetrameric symmetry of K\textsuperscript{+} channels and the simplicity of 4 transitions without memory that describe $I_{\rm Ks}$ activation. This detailed description of activation, in particular the delay before activation, is a channel feature that has not previously been incorporated into an $I_{\rm Ks}$ model.

Model parameters were determined with the use of non-linear optimization. Although we incorporated a large set of experiments, there may be different parameters that also reproduce channel properties. Our conclusions depend on the participation of numerous closed states in $I_{\rm Ks}$ activation. The necessity of these states for generating a delay before activation,\textsuperscript{14,19} in particular the delay before activation, is a channel feature that has not previously been incorporated into an $I_{\rm Ks}$ model.

Future work should incorporate \beta-adrenergic modulation of $I_{\rm Ks}$ and study its effects on whole-cell electrophysiological function. To accomplish this, a \beta-adrenergic model that simulates its effects on many cellular processes is necessary; these processes include sarcoplasmic reticulum calcium handling, $I_{\rm CaL}$, the transient outward Cl\textsuperscript{-} current ($I_{\rm o2}$), $I_{\rm Na}$, $I_{\rm ScCl}$, $I_{\rm Ks}$, and $I_{\rm K1}$. To date there have been attempts to create such a model,\textsuperscript{21,22} but a sufficiently complete model that allows accurate study of AP dynamics in the context of \beta-adrenergic stimulation awaits development.

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