New Aspects of HERG K⁺ Channel Function Depending upon Cardiac Spatial Heterogeneity

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Abstract
HERG K⁺ channel, the genetic counterpart of rapid delayed rectifier K⁺ current in cardiac cells, is responsible for many cases of inherited and drug-induced long QT syndromes. HERG has unusual biophysical properties distinct from those of other K⁺ channels. While the conventional pulse protocols in patch-clamp studies have helped us elucidate these properties, their limitations in assessing HERG function have also been progressively noticed. We employed AP-clamp techniques using physiological action potential waveforms recorded from various regions of canine heart to study HERG function in HEK293 cells and identified several novel aspects of HERG function. We showed that under AP-clamp IHERG increased gradually with membrane repolarization, peaked at potentials around 20–30 mV more negative than that revealed by pulse protocols and at action potential duration (APD) to 60%-70% full repolarization, and fell rapidly at the terminal phase of repolarization. We found that the rising phase of IHERG was conferred by removal of inactivation and the decaying phase resulted from a fall in driving force, which were all determined by the rate of membrane repolarization. We identified regional heterogeneity and transmural gradient of IHERG when quantified with the area covered by IHERG trace. In addition, we observed regional and transmural differences of IHERG in response to dofetilide blockade. Finally, we characterized the influence of HERG function by selective inhibition of other ion currents. Based on our results, we conclude that the distinct biophysical properties of HERG reported by AP-clamp confer its unique function in cardiac repolarization thereby in antiarrhythmia and arrhythmogenesis.

Introduction
Cardiac function, in particular the cardiac electrical activity, is controlled by an array of ion channels that activate in a well-organized manner with a delicate balance between inward and outward ion movements. Upon receiving an incoming impulse, cardiac cells excite with a rapid membrane depolarization followed by a relatively slow repolarization process. Repolarization abnormalities, either excessive slowing or accelerating of the rate of repolarization, can cause cardiac electrical perturbations or arrhythmias. The rate of repolarization is determined by a team effort of several ion currents, of which the rapid delayed rectifier K⁺ current (IKr) is a crucial player, particularly to the plateau phase of an action potential [1–5]. The major molecular component of native IKr has been identified as the human ether-a-go-go-related gene (HERG) which when expressed in heterologous systems generates IKr-like current [4]. Impairment of IKr/HERG can cause excessive prolongation of action potential duration (APD). Indeed, HERG has been implicated as a molecular target for mutations leading to induction of chromosome-7 linked type 2 familial long QT syndrome and as a pharmacological target for drug actions that induce acquired long QT syndrome, a lethal ventricular tachyarrhythmia often leading to sudden cardiac death [4].

HERG possesses unique biophysical characteristics and pharmacological properties distinct from other K⁺ channels. During membrane depolarization, HERG K⁺ channel simultaneously activates and inactivates; but its inactivation kinetics is faster than its activation kinetics, with inactivation time constants (in the order of ms) some 1–2 orders of magnitude smaller than the activation time constants (in a range up to hundreds of ms) at the same potential. Upon membrane repolarization or hyperpolarization, HERG channel rapidly reactivates from inactivated state and slowly deactivates; the reactivation kinetics is similar to the inactivation time course whereas the deactivation process is approximately two orders of magnitude slower than the inactivation, at the same membrane potential. These biophysical characteristics confer HERG a strong inward rectification and a bell-shaped current-voltage relationship (I–V relationship). Presumably due to its wide-opened pore architecture, HERG channel is susceptible to blockade by a variety of drugs and compounds of various categories with heterogeneous structures [5]. This high pharmacological sensitivity renders HERG the most frequent cause of drug-induced long QT syndrome. In addition, HERG might also be involved in pathological QT prolongation. Heart failure is electrophysiologically characterized by abnormal QT prolongation and early afterdepolarization (EAD) [6]. Indeed, depression of IKr has been documented in failing hearts of both animal models [7,8] and theoretical simulation [9]. Tumor necrosis factor-α (TNF-α), a critical factor causing heart failure and when overexpressed prolonging APD and promoting ventricular arrhythmias, suppresses HERG function [10]. Recent studies further suggest that decreasing IKr increases rate-related APD...
alternants that can dispose to ventricular tachyarrhythmias [11,12]. On the other hand, enhancement of HERG function might also be proarrhythmic. A study demonstrated that IKr-like current is increased in cells from acute myocardial infarction that is associated with arrhythmias [13,14]. And it has been shown that HERG function is enhanced by phospholipid metabolites that are known to accumulate during the early phase of acute myocardial ischemia and to cause ischemic arrhythmias by shortening APD [10,15–17]. These data indicate that IKr/HERG plays an important role in determining propensity of arrhythmias. Understanding HERG function is therefore crucial to understanding mechanisms of arrhythmias of various types under different pathological conditions or induced by drug toxicity.

A cardiac action potential (AP) is typically divided into phase 0 depolarization, phase 1 repolarization, phase 2 plateau, and phase 3 repolarization, with magnitude, length and rate of these phases varying in cells from different regions of the heart. The key feature of cardiac AP is its relatively long repolarization process, ranging from decades of to hundreds of milliseconds depending on species and regions from which the cells are investigated, compared to its rapid and brief phase 0 depolarization (within 2 ms). This implies that during each excitation a cardiac cell spends a majority of time in the repolarization process after its peak depolarization. Another important note about the cardiac APs is the variability of AP waveforms; the regional differences (such as sinus-atrial nodal (SAN), atrial and ventricular cells) and transmural heterogeneity (such as endocardium, midmyocardium and epicardium of ventricular walls) [18–20]. Notably, given the unique features of voltage- and time-dependence of HERG channel and the characteristics of cardiac APs, HERG channel activities are expected to differ during membrane repolarization from during depolarization, and HERG function assessed with repolarizing waveforms (or AP configurations) should be of more physiological relevance. However, HERG function has been studied primarily by conventional voltage step protocols with long rectangular depolarizing pulses of usually ≥2 s in duration—non-physiological voltage waveforms. This view raises a serious doubt about the fidelity of depolarizing pulses to reflect the true physiological function and pharmacological sensitivity of HERG K+ channels. Some efforts have been made in previous studies to address this apparent problem by using either authentic AP waveforms or simulated AP configurations in assessing HERG function [11,12,21–23]. These studies indeed revealed some important features of HERG channels, which are otherwise masked by conventional square pulse protocols. Nonetheless, precise understanding of HERG activity under more physiological conditions during an AP is still lacking. Specifically, several unanswered questions await more detailed investigation. First, what are the biophysical attributes that determine HERG function during an AP in cardiac cells? Or conversely, how do changes of HERG function influence cardiac repolarization? Second, whether there exist regional differences in terms of HERG function and if yes how do these differences contribute to the highly organized cardiac repolarization? Finally, how does IKr/HERG change in response to inhibition of ion currents that activate before IKr/HERG does? The present study was designed to delineate these points with the use of AP-clamp techniques.

**Methods**

**Cell Culture**

HEK293 cells stably expressing HERG were grown in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% heat-inactivated fetal bovine serum, 200 μM G418, 100 units/ml penicillin, and 100 μg/ml streptomycin. For electrophysiological study, cells subcultured to ~85% confluency were harvested by trypsinization and stored in the Tyrode solution containing 0.5% bovine serum albumin at 4°C [24]. Cells were studied within 10 hours of harvest.

**Myocyte Isolation**

For action potential clamp (AP-clamp) experiments, APs were obtained from myocytes isolated from different regions of canine hearts. The procedure for canine myocyte isolation was essentially the same as described in detail elsewhere [25]. Briefly, adult male mongrel dogs were anesthetized with morphine (2 mg/kg SC) and α-chloralose (120 mg/kg IV load, 29.25 mg/kg/h infusion), and a median sternotomy was performed. The hearts were removed and mounted to a Langendorff perfusion system. The preparation was perfused with Ca2+-containing Tyrode solution at 37°C until the effluent was clear of blood, and the perfusate was then switched to Ca2+-free Tyrode solution for 20 min at a constant rate of 25 ml/min, followed by perfusion with the same solution containing collagenase and 0.1% BSA for 35 min. The right atrium was dissected out in Krafthruhe (KB) medium and cells were dispersed by agitation. Dissection of different layers of left ventricular wall was carried out according to the method established by Liu et al [26]. Briefly, after perfusion, thin slices of tissues were dissected from epicardium (<1.5 mm from epicardial surface), M region (3 to 5 mm from epicardial surface), and endocardium (<2 mm from the endocardial surface). Shavings were made parallel to the surface of the left ventricular free wall midway along the apicobasal axis. The SAN region was dissected out and cut into small pieces in the enzyme solution for continuous digestion for another 20 min. Cells were then stored in KB medium for later use. A total of six adult male mongrel dogs (22–27 kg) were used in this study. The protocols for the use of dogs in this study were approved by the Animal Use and Care Committee of the Minhang Central Hospital.

**Whole-Cell Patch-Clamp Recording**

Patch-clamp recording of HERG K+ currents (IHERG) was conducted with the whole-cell voltage-clamp methods in the voltage-clamp mode, and single cell action potential was recorded in the current-clamp mode, using an Axopatch 200B amplifier (Axon Instruments, Burlingame, CA). Borosilicate glass electrodes (1 mm) had tip resistances of 1 to 3 MΩ when filled with pipette solution containing (mM) 130 KCl, 1 MgCl2, 5 Mg-ATP, 10 EGTA, and 10 HEPES (pH 7.3). Junction potentials were zeroed before formation of the membrane-pipette seal in the extracellular (Tyrode) solution containing (mM) 136 NaCl, 5.4 KCl, 1 CaCl2, 1 MgCl2, 10 glucose, and 10 HEPES (pH 7.4). Single cell action potentials (APs) in myocytes isolated from various regions of canine hearts were revoked by 10 consecutive stimuli of 3-ms duration and twice threshold strength at a frequency of 1 Hz (or a cycle length of 1 s). The APs recorded by the 10th pulse reached the steady state and were converted to pCLAMP 6 format for use as voltage-clamp waveforms using the DacFile command in pCLAMP 6. For current recordings in HERG-expressing HEK293 cells, the capacitance and series resistance were electrically compensated to minimize the duration of the capacitative surge on the current recording and the voltage drop across the clamped cell membrane. Cells with changing leak current (indicated by >10 pA changes in holding current at 250 mV) were rejected. Experiments were conducted at 36±1°C. All chemicals were purchased from Sigma Chemical Co.
Data Analysis

Group data are expressed as mean ± S.E. Statistical comparisons (performed using ANOVA followed by Dunnett’s method) were carried out, and paired or unpaired t-test was used, as appropriate, for single comparisons, using Microsoft Excel. A two-tailed p<0.05 was taken to indicate a statistically significant difference. Nonlinear least square curve fitting was performed with CLAMPFIT in pCLAMP 8.0 or GraphPad Prism and integration was performed to calculate the area under \( I_{\text{HERG}} \) trace using GraphPad Prism.

Results

Characteristics of Action Potentials from Various Regions of the Heart

Action potentials (APs) were recorded in cardiomyocytes isolated from various regions of canine hearts, including endocardium, midmyocardium and epicardium of left ventricular wall, right atrium and SAN. APs from different regions had different morphology (Fig. 1) and characteristics (Table 1).

The AP from epicardium (Epi-AP) was characterized by prominent spike-and-dome morphology, with an initial rapid and pronounced repolarization followed by a secondary depolarization. Midmyocardial AP (Mid-AP) demonstrated a moderate phase 1 spike and a long flat phase 2 plateau. The APs from three layers of left ventricular wall all had a rapid and regenerative phase 4 repolarization. By comparison, typical atrial AP (A-AP) was characterized by a short plateau phase and a slow final repolarization, approximating triangular configuration. The AP from sinus-atrial nodal cells (SAN-AP) typically displayed a slow spontaneous depolarization followed by a rapid depolarization and short action potential duration (APD). To quantitatively characterize the APs, several parameters important for analyzing HERG function were measured and are summarized in Table 1. \( \text{APD}_{10} \), \( \text{APD}_{20} \), \( \text{APD}_{50} \), \( \text{APD}_{60} \) and \( \text{APD}_{90} \) were determined for APD to 10%, 20%, 50%, 60% and 90% of repolarization, respectively.

| Table 1. Characteristics of Action Potentials Recorded From Various Regions of Canine Heart and Used as Command Waveforms for AP-Clamp. |
|---------------------------------------------------------------|
|                  | Endo-AP | Mid-AP | Epi-AP | A-AP | SAN-AP |
| \( \text{APD}_{10} \) (mV - ms) | 29 — 15 | 34 — 13 | 31 — 13 | 27 — 17 | 11 — 44 |
| \( \text{APD}_{20} \) (mV - ms) | 17—65 | 21—155 | 19—32 | 16—30 | 3—58 |
| \( \text{APD}_{50} \) (mV - ms) | -20—197 | -17—366 | -18—230 | -19 | 82 | -21; 72 |
| \( \text{APD}_{60} \) (mV - ms) | -32—219 | -30—378 | -31—245 | -30—96 | -29—75 |
| \( \text{APD}_{90} \) (mV - ms) | -44—232 | -42—384 | -43—249 | -46—135 | -41—80 |
| RR2 (mV/sec) | 39 | 22 | 36 | 45 | 40 |
| RR3 (mV/sec) | 72 | 80 | 75 | 35 | 65 |
| RR4 (mV/sec) | 72 | 80 | 75 | 35 | 65 |
| SDR (mV/sec) | N/A | N/A | -65 | N/A | N/A |

Figure 1. Action potential (AP) waveforms recorded from myocytes isolated from various regions of the canine heart. These AP waveforms were used for the AP-clamp studies of HERG K⁺ channel function in HERG-expressing HEK293 cells. The quantitative characteristics of these APs are summarized in Table 1. doi:10.1371/journal.pone.0072181.g001
and the voltages at the respective durations. AP amplitude (APA) and diastolic membrane potential (DMP) were measured as total amplitude of APs and voltage at diastolic interval, respectively. The rates of membrane repolarization for phase 2, phase 3 and phase 4 (RR2, RR3, and RR4, respectively) were determined by linear regressions to the steepest portion of the corresponding repolarizing phases, and the rate of secondary depolarization (SDR) was also calculated.

The rate of initial repolarization (APD0) was similar between various layers of ventricle and atrium, but was considerably slower in SAN. The transmural difference became prominent from the subsequent phases of repolarization. For instance, APD50 of Mid-AP was around 2.5-fold and 3-fold longer than Endo-AP and Epi-AP, respectively, in the order of Mid-AP > Endo-AP > SAN-AP > Epi-AP = A-AP. The length of Epi-AP began to exceed that of Endo-AP from phase 2 repolarization; the order of the length of APD50, APD60, and APD90 was Mid-AP < A-AP < SAN-AP. RR1, APA and DMP were in the same range for V-APs and A-AP, and lower for SAN-AP.

Characteristics of HERG K⁺ channel kinetics by conventional pulse protocols

To aid understanding IHERG function and biophysics with AP-clamp, comparison with the conventional pulse protocol was made. Depolarizing voltage steps from −60 mV to +40 mV elicited delayed rectifier outward currents with characteristic inward rectification that was manifested at stronger depolarization. Upon repolarization back to −50 mV, slowly decaying tail currents were recorded. The inward rectification resulted in a bell-shaped I-V relationship with IHERG peaking at −15 mV (Fig. 2A).

**Steady-state voltage-dependent activation.** The steady-state voltage-dependent activation was determined by tail currents elicited by the voltage protocol shown in the inset of Fig. 2B. The half-maximal inactivation voltage (V1/2) and the slope factor (k) determined by the Boltzmann distribution were −30±2 mV and −5 mV, respectively.

**Steady-state voltage-dependent inactivation.** The steady-state voltage-dependent inactivation was assessed by the voltage protocol shown in the inset of Fig. 2C. IHERG was first inactivated by a 2-s depolarizing step to +40 mV and then reactivated to various extents during the hyperpolarizing pulses to various potentials of a 10-ms duration that allowed for full reactivation with minimal deactivation, and the decaying outward currents induced by the subsequent 50 ms depolarizing pulse to +20 mV represent IHERG inactivation. The half-maximal inactivation voltage (V1/2) and the slope factor (k) determined by the Boltzmann distribution are −55±5 mV and −18 mV, respectively.

**Activation kinetics.** The apparent activation time constants (τact) of HERG channels were determined by the single exponential fit to the step IHERG during depolarizing pulse to 0 mV. The values ranged from 330 ms to 50 ms between voltage steps of −20 mV and +20 mV (Fig. 2B).

**Deactivation kinetics.** To evaluate the voltage-dependence of the apparent deactivation time constants (τdeact) of HERG channels, the voltage protocol as shown in Fig. 2B was applied. The decaying phase of the tail IHERG elicited at various hyperpolarizing/repolarizing steps (from −120 mV to −50 mV), following a depolarizing prepulse to +40 mV, were fit by the single exponential to obtain τdeact. It is important to note that apparent deactivation process was absent at potentials positive to −50 mV, and minor and slow deactivation was seen at potentials from −50 mV to −70 mV with time constants of 3.8 s to 1.4 s.

**Inactivation kinetics.** The inactivation kinetics (τinact) was analyzed with the voltage protocol shown in the inset of Fig. 2C. The decaying tail currents recorded upon depolarization to various potentials ranging from −80 mV to +40 mV to inactivate HERG channels, preceded by a 10-ms hyperpolarizing prepulse to −120 mV to let channels fully recover, were fit to the monoexponential function and the values of τinact ranged from 0.5 ms to 1.3 ms between −80 mV and +40 mV.

**Reactivation kinetics.** The reactivation time course (τreact) was determined with the same voltage protocol as that for steady-state inactivation. The monoexponential fit was applied to the activating phase of the tail IHERG evoked at various hyperpolarizing/repolarizing steps following the inactivation steps at +40 mV. The reactivation time constants (τreact) range from 8.2 ms to 1.2 ms for voltages from −120 mV to −40 mV.

These results indicate that (1) with respect to voltage, HERG channel inactivated from more negative potentials than it activates (−55 mV vs. −30 mV) and (2) with respect to time, the time courses of inactivation and reactivation from inactivation were comparable, and were 1–2 orders of magnitude faster than the activation and 3 orders of magnitude faster than the deactivation processes within the physiological voltages spanned by an AP.

**General Characteristics of IHERG Assessed by Action Potential Waveforms from Different Areas of the Heart**

On the basis of our data with conventional square pulse protocols, we predicted that the activation of HERG is rapid during an AP. However, due to rapid inactivation relatively little current flows until the potential becomes less positive than 0 mV. The removal of inactivation then allows more current to flow, giving rise to the distinct pattern of IHERG changes during the AP. IHERG elicited by AP-clamp could be roughly divided into 2–3 rising phases and 2 decaying phases. The initial rising phase was rapid and corresponded to the phase 0 upstroke of AP waveform. This was followed by a slowly rising phase corresponding to the phase 2 plateau and a subsequent fast rising phase during phase 3 repolarization, with ventricular AP waveforms (V-APs). With atrial AP (A-AP) and SAN-AP waveforms, the second slowly rising phase was absent presumably because of the short phase 2 plateau duration. After reaching the peak, IHERG rapidly decayed during the final repolarization. Subsequent to the fast decay was a slowly decaying phase occurring after complete repolarization, equivalent to the tail IHERG recorded upon repolarization steps (Fig. 3). Note that there was an inward tail at the end of the trace after complete repolarization of V-APs. This was likely due to the fact that the resting membrane potential of our cells was around −74 mV and repolarizing to −80 mV with V-APs generated inward current, whereas repolarizing to −70 mV with A-AP or to −55 mV with SAN-AP produced outward tail currents.

Detailed analysis of AP-clamp revealed several novel aspects of IHERG important to our understanding of HERG function and biophysics, which would otherwise be masked by the conventional step protocols. First, although the I-V relationship of AP-clamp IHERG was bell-shaped, similar to the step IHERG, the potentials at which IHERG peaked were around 20 mV more negative with AP-clamp than with conventional step clamp. Specifically, IHERG peaked at −10 mV with pulse protocols, but at −30 to −40 mV with AP waveforms (Fig. 3 and Table 2). With respect to APD, IHERG peaked at the time when membrane potential reached 60–70% full repolarization, that is, APD50 and APD70 (Fig. 3 and Table 2). For more straightforward view of these relationships, direct comparison between AP configuration and IHERG waveform is depicted in Fig. 4A. The vertical dash lines indicate the position where IHERG peaks in relation to APD. Note that the amplitude and duration scales are the same for all recordings in Fig. 4A and the IHERG of Endo-AP peaked to the highest level, while the IHERG.
of SAN-AP peaked to the lowest level, relative to the respective AP amplitudes (horizontal dash lines).

Second, to quantitatively describe the time-dependent changes of IHERG during an AP waveform, we calculated the slope (or the rates) of various rising and decaying phases by linear regression to the data points spanning the steepest portion of the given phases (Fig. 4B). The corresponding phases of APD were also analyzed for the rates of repolarization by the same methods (Fig. 4B). The resulting slopes were used to measure the rates of ascending and descending phases of IHERG. Comparison between the rates of IHERG changes and the rates of various phases of AP waveforms revealed that the rates of IHERG rising phases and decaying phase depended upon the rates of corresponding phases of membrane repolarization. That is, the greater the rate of repolarization (the steeper of the phase) of an AP was, the faster the rising (or decaying) phase of IHERG was. This point was further reinforced...
by the fact that when the AP and the corresponding \( I_{\text{HERG}} \) trace were superimposed, the repolarizing phases of the AP and the rising phases of the \( I_{\text{HERG}} \) had a nearly perfect mirror-image relationship (Fig. 4A).

Third, both the voltage-dependence and time-dependence of changes of the instantaneous conductance of HERG channels (\( G \)) were analyzed according to the equation \( G = I/V \) (where \( I \) represents a given \( I_{\text{HERG}} \) amplitude at the corresponding voltage (\( V \)) during AP-clamp). As shown in Fig. 5, the \( G-V \) relationship with V-APs and A-AP was sigmoidal and can be fit by the Boltzmann function, and the \( G-V \) relationship with SAN-AP waveform was exponential. The \( G-V \) relationship could be viewed as an instantaneous inactivation curve with conductance or channel availability decreasing as membrane depolarizes or as an instantaneous reactivation curve with conductance or channel availability increasing as membrane repolarizes; with respect to AP waveform, the latter might be more informative. The resulting instantaneous \( V_{1/2} \) values for Epi-AP (−51 mV) and Mid-AP (−51 mV) were similar to that determined by pulse protocols and for Endo-AP (−37 mV) and A-AP (−38 mV) the values corresponded to more positive potentials. By comparison, the \( k \) values for Endo-AP (−18 mV) and A-AP (−14 mV) were close to that with pulse protocols and those for Epi-AP (−28 mV) and Mid-AP (−26 mV) were much shallower. When \( G_m \) was expressed as a function of time (\( T \)) or duration, we obtained the instantaneous \( G-T \) relationship of HERG channels. The \( G-T \) relationship showed exponential increases with F-APs while sigmoidal increases with A-AP and SAN-AP waveforms.

Finally, as already described above, \( I_{\text{HERG}} \) peaked at around APD_{60} and APD_{70} with respect to APD and at between −30 mV and −40 mV with respect to membrane potential. Evidently, the rising phase of \( I_{\text{HERG}} \) was mainly due to reactivation of HERG channels during repolarization from inactivation rendered upon initial depolarization. However, it was unclear what determines the decaying phases of \( I_{\text{HERG}} \) during AP-clamp. One possibility is deactivation of HERG channels as commonly believed [21,23,27]. Nonetheless, close inspection of deactivation process with step protocol revealed that the deactivation time course was slow and virtually absent at potentials positive to −50 mV (Fig. 2B), implying that at the voltages where \( I_{\text{HERG}} \) started to decline, no channel deactivation occurred. It is therefore unlikely that the decaying of \( I_{\text{HERG}} \) during an AP is ascribed to deactivation process. To resolve the issue, we calculated the instantaneous driving forces (DF) during an AP waveform by subtracting the voltages at given time points from the reversal potential of \( I_{\text{HERG}} \) in a given cell. DF and \( G \) were then normalized to their own maximum values. The predicted \( I_{\text{HERG}} \) amplitude was determined by multiplying DF by \( G \) (\( I = G \times DF \)). As shown in Fig. 6, the values were maximal at around −30 mV in voltage and APD_{60} in duration for V-APs and −40 mV and APD_{70} for A-AP and SAN-AP. Noticeably, at durations after APD_{60} and APD_{70}, \( G \) continually increased while DF began to reduce. The data suggest

**Figure 3.** \( I_{\text{HERG}} \) waveforms and I-V relationships measured by AP-clamp. Left panels: Typical examples of current waveforms recorded by AP-clamp; Right panels: I-V relationships averaged from at least 15 cells. Dash lines indicate the voltages at which \( I_{\text{HERG}} \) peaks (\( V_{\text{Peak}} \)). The mean \( V_{\text{Peak}} \) was averaged from 15 cells for each group. *\( p<0.05 \) vs. Endo.

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Table 2. Characteristics of $I_{\text{HERG}}$ Reported by AP-Clamp.

|                    | Endo-AP | Mid-AP | Epi-AP | A-AP | SAN-AP |
|--------------------|---------|--------|--------|------|--------|
| $V_{\text{Peak}}$ (mV) | $-34 \pm 1$ | $-30 \pm 2$ | $-32 \pm 1$ | $-41 \pm 1$ | $-39 \pm 0$ |
| APD$_{\text{Peak}}$ (% Repolarization) | $62 \pm 1$ (APD$_{50}$) | $60 \pm 2$ (APD$_{50}$) | $61 \pm 1$ (APD$_{50}$) | $70 \pm 1$ (APD$_{50}$) | $73 \pm 1$ (APD$_{50}$) |
| $A$ (pA/pF*s×1000) | $-17 \pm 1$ | $-15 \pm 1$ | $-11 \pm 1$ | $-11 \pm 1$ | $-6 \pm 1$ |
| $V_{1/2, \text{IT}}/k$ (mV) | $-37/19$ | $-50/28$ | $-51/26$ | $-38/14$ | N/A |
| RP2 (pA/ms) | $21 \pm 1$ | $8 \pm 0$ | $13 \pm 1$ | $35 \pm 1$ | $22 \pm 2$ |
| RP3 (pA/ms) | $45 \pm 3$ | $50 \pm 3$ | $50 \pm 3$ | $55 \pm 4$ | $26 \pm 2$ |
| DP1 (pA/ms) | $-80 \pm 4$ | $-85 \pm 5$ | $-32 \pm 1$ | $-50 \pm 3$ | $-45 \pm 5$ |
| DP2 (pA/ms) | N/A | N/A | N/A | $-12 \pm 1$ | N/A |

$V_{\text{Peak}}$: the voltage at which $I_{\text{HERG}}$ peaks; APD$_{\text{Peak}}$: % repolarization at which $I_{\text{HERG}}$ peaks; hence, the value 62% in the table can be expressed as APD$_{62}$; $A$: area under $I_{\text{HERG}}$ trace, a measure of the volume of K$^{+}$ passing through HERG channel during an AP; $V_{1/2, \text{IT}}$: voltage for half maximum inactivation calculated from the Boltzmann fit to the instantaneous inactivation curve; $k$: slope factor from the Boltzmann fit for instantaneous conductance curve; RP2 and RP3: rising phase 2 and 3 of $I_{\text{HERG}}$ trace, respectively; DP1 and DP2: decaying phase 1 and 2 of $I_{\text{HERG}}$ trace, respectively. Positive sign indicate ascending and negative sign descending directions.

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Figure 4. Direct comparison between AP of various regions and the corresponding $I_{\text{HERG}}$ waveforms. A: Superimposed AP (dotted line) and $I_{\text{HERG}}$ (solid line) traces. Vertical dash lines indicate AP at which $I_{\text{HERG}}$ peaks (APD$_{\text{Peak}}$). Note that the horizontal dash lines pass through the converging point between AP and $I_{\text{HERG}}$ and form the boundary for the mirror-image relationship between AP and $I_{\text{HERG}}$. The mean data of APD$_{\text{Peak}}$ were averaged from 15 cells for each group. The amplitude and duration scales are the same for all recordings. “$p < 0.05$ vs. Endo.” B: AP and $I_{\text{HERG}}$ traces showing the measurements of slopes of various phases by linear regression indicated by the straight lines, as a measure of rate of repolarization for AP and rising/decaying phase for $I_{\text{HERG}}$: RR2, RR3 and RR4, repolarization rate corresponding to the phases of APD$_{50}$, APD$_{60}$, and APD$_{90}$, respectively. RP2 and RP3: rising phase 2 and 3 of $I_{\text{HERG}}$ trace, respectively; DP1 and DP2: decaying phase 1 and 2 of $I_{\text{HERG}}$ trace, respectively.

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Figure 5. Time- and voltage-dependence of the instantaneous conductance of HERG K⁺ channels reported by AP-clamp. The instantaneous conductance (G) was calculated by dividing the current amplitude by the membrane potential of an AP. Left panels: G as a function of time (APD); Right panel: G as a function of membrane potential. The latter is considered as instantaneous inactivation. The lines represent the Boltzmann fit to the experimental data points.

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that the rapid decaying phase of $I_{\text{HERG}}$ with AP-clamp was a consequence of falling of DF.

Spatial Heterogeneity of $I_{\text{HERG}}$ Revealed by AP-Clamp

Comparison of AP-clamp $I_{\text{HERG}}$ revealed some important regional differences of HERG function in terms of its voltage-dependent and time-dependent properties.

First, $I_{\text{HERG}}$ with V-APs from three different layers all peaked at approximately $230 \text{ mV}$ and with A-AP and SAN-AP at around $240 \text{ mV}$ (Fig. 3 and Table 2).

Second, $I_{\text{HERG}}$ elicited with V-APs peaked at duration equivalent to APD$_{60}$, despite the transmural differences in the length of APD$_{60}$. By comparison, $I_{\text{HERG}}$ recorded with A-AP or SAN-AP peaked at duration to APD$_{70}$ (Fig. 4 and Table 2). These corresponded to 219 ms, 378 ms and 245 ms for APD$_{60}$ of Endo-AP, Mid-AP and Epi-AP, respectively, and 135 and 80 ms for APD$_{70}$ of A-AP and SAN-AP, respectively.

To better quantify the size of $I_{\text{HERG}}$, we calculated the area covered by $I_{\text{HERG}}$ trace elicited by an AP using the integration function of Graphpad Prism software and setting the baseline at the zero current level; such analysis gives more accurate measure of $K^{+}$ flux carried by HERG channels during an AP. $I_{\text{HERG}}$ determined in this way demonstrated spatial differences and as expected, the area was in general greater in the region with longer APD. For instance, the length of APD was in the order of V-APD$_{60}$, A-APD$_{60}$, and correspondingly, the size of $I_{\text{HERG}}$ area was also in the same order (Fig. 7A).

Transmural Heterogeneity of $I_{\text{HERG}}$ Revealed by AP-Clamp

Unexpectedly, when comparing $I_{\text{HERG}}$ elicited with AP waveforms across the left ventricular wall, the area, in the order of Endo-AP>Mid-AP>Epi-AP (Fig. 7A), did not correlate with the length of APD alone. Apparently, other factors in addition to APD also play an important role in determining the size of $I_{\text{HERG}}$ area. As already mentioned above, HERG reactivation or reopening depended upon the rate of membrane repolarization and DF determines the size of $I_{\text{HERG}}$. This implies that the rate of repolarization (a measure of HERG conductance $G$) and DF during various phases of an AP should be taken into consideration for understanding the spatial heterogeneity of $I_{\text{HERG}}$ area. We then formulated the following equations to predict relative size of $I_{\text{HERG}}$ area of different regions:

$$A = RR2 \times APD50 \times DF2 + RR3 \times APD60 - 50 \times DF2 + RR4 \times APD90 - 60 \times DF4$$  \hspace{1cm} (1)

$$A = RR2 \times APD50 - 20 \times DF2 + RR3 \times APD60 - 50 \times DF2 + RR4 \times APD90 - 60 \times DF4 - SDR2 \times APD20 \times DF1$$  \hspace{1cm} (2)

$$A = RR2 \times APD50 \times DF2 + RR3 \times APD70 - 50 \times DF2 + RR4 \times APD90 - 70 \times DF4$$  \hspace{1cm} (3)

where $A$ represents the area of $I_{\text{HERG}}$, RR the rate of repolarization of the given phase of an AP as specified by the number, and DF the mean driving force of the given phase of an AP as specified by the number. APD$_{50}$ approximates the length of the plateau phase, APD$_{60-50}$ (APD$_{60}$-APD$_{50}$) the duration of transition from phase 2 to phase 3 repolarization, and APD$_{90-60}$ (APD$_{90}$-APD$_{60}$) the duration phase 3 repolarization. SDR$_{2}$ indicates the rate of secondary depolarization and APD$_{20}$ approximates the duration of the secondary depolarization, which were seen with Epi-AP because of the spike-and-dome morphol-
Ion currents activate in a certain sequence during an action potential. The one that activates in earlier phases may affect activation of the ones that activate in later phases by setting the voltage and time preconditions for the channel state. For example, the transient outward K$^+$ current ($I_{\text{to}}$) activates in the early phase of an AP and alteration of $I_{\text{to}}$ may change the function of the subsequently activated currents such as $I_{Kr}$, by altering AP morphology (voltage and duration). In a previous study, the authors showed that nicotine selectively inhibits $I_{\text{to}}$ at low concentrations without affecting other K$^+$ currents [28].

To investigate the potential influence of $I_{\text{to}}$ alteration on $I_{Kr}$, we recorded APs in the presence of nicotine (500 nM) to inhibit $I_{\text{to}}$ in canine atrial myocytes and these APs have elevated plateau level and markedly shortened APD. The ultrarapid delayed rectifier K$^+$ current ($I_{\text{Kur}}$) is an atrium-specific and activates in the early phase of AP with partial inactivation, contributing to the plateau repolarization of atrial myocytes [25,29]. It is highly sensitive to 4-aminopyridine (4-AP) block, with micromolar concentrations of 4-AP nearly abolishing $I_{\text{Kur}}$ without significantly affecting other ion currents and blockade of $I_{\text{Kur}}$ produces some 50–60% APD lengthening. Here the APs recorded from canine right atrial cells in the presence of 100 µM 4-AP to block $I_{\text{Kur}}$ have elevated plateau level and prolonged APD. The characteristics of L-type Ca$^{2+}$ current ($I_{\text{CaL}}$) blockade by 10 µM nisodipine is the loss of plateau phase and shortening of APD, particularly the early phase repolarization (Fig. 9). The $I_{\text{HERG}}$ elicited by the AP waveforms recorded under these different conditions demonstrated similar but different characteristics, as shown in Fig. 9. $I_{\text{HERG}}$ peaked at 110 ms under control conditions and when $I_{\text{to}}$ was inhibited (labeled $I_{\text{to}^-}$). When $I_{\text{Kur}}$ was blocked (labeled $I_{\text{Kur}^-}$), the peak of $I_{\text{HERG}}$ shifted to 140 ms, whereas when $I_{\text{CaL}}$ was blocked (labeled $I_{\text{CaL}^-}$) $I_{\text{HERG}}$ peaked sooner at 75 ms. In respect with voltage, the maximum $I_{\text{HERG}}$ occurred at around −40 to −43 mV under control conditions or with $I_{\text{Kur}^-}$ or $I_{\text{CaL}^-}$, and it was some 10 mV more negative shift when $I_{\text{to}}$ was inhibited. The most notable difference was the size of $I_{\text{HERG}}$ area. The characteristics of the atrial APs used for AP-clamp and the corresponding $I_{\text{HERG}}$ in the presence of varying K$^+$ channel blockers to inhibit $I_{\text{to}}$, $I_{\text{Kur}}$ or $I_{\text{CaL}}$, separately, are summarized in Table 3.

Regional Differences of $I_{\text{HERG}}$ in Response to Dofetilide Blockade Revealed by AP-Clamp

Since the effect of $I_{Kr}$/I$\text{HERG}$ blockers is state-dependent with most of them generally acting on the open channels and since there is regional difference of APs in terms of the repolarization rate thereby the rate of HERG reopening, we reasoned that the effects of $I_{Kr}$/I$\text{HERG}$ blockers are different in cells from different regions of the heart. To test this notion, we went on to study the effects of dofetilide on $I_{\text{HERG}}$ using AP-clamp with AP waveforms collected from various regions. As illustrated in Fig. 9, 10 nM dofetilide which produced approximately 50±3% inhibition of $I_{\text{HERG}}$ area with Endo-AP reduced the current by 27±3%, 22±1%, 21±6%, and 30±3% with Mid-AP, Epi-AP, A-AP and SAN-AP, respectively (Fig. 9A and 9B). When the data were normalized to Epi-AP, it was clear that the degree of blockade of $I_{\text{HERG}}$ with A-AP was around 4-times greater than with Epi-AP and about 2–3-times greater than in APs of other regions (Fig. 9C). Also important is that there were transmural differences of $I_{\text{HERG}}$ in response to dofetilide block. For example, the blockade of $I_{\text{HERG}}$ with Endo-AP was 2.5-fold greater than with Epi-AP and 2-fold greater than with Mid-AP (Fig. 9C). Noticeably, these differential effects of dofetilide caused a remarkable change in the pattern of transmural difference of $I_{\text{HERG}}$; the normal $I_{\text{HERG}}$ gradient was changed to Mid-AP $>$ Epi-AP $>$ Endo-AP by dofetilide.

Figure 7. Spatial and transmural heterogeneity of HERG K$^+$ channel function. A. Experimental data of the area under $I_{\text{HERG}}$ trace, a quantitative index of the size of current flow through HERG channels during action potentials. *p<0.05 vs. Endo; +p<0.05 F-test (ANOVA) for transmural difference. B. Theoretical (Predicted) data of the area under $I_{\text{HERG}}$ trace. C. Correlation between experimental data and predicted data. The solid line represents the linear regression to the data points and the dotted lines indicate 95% confidence. doi:10.1371/journal.pone.0072181.g007

Alterations of $I_{\text{HERG}}$ in Response to Blockade of $I_{\text{to}}$, $I_{\text{Kur}}$, or $I_{\text{CaL}}$

$I_{\text{HERG}}$ properties of HERG channels reported by AP-clamp

- Equation 1 was used for Endo-AP and Mid-AP, equation 2 for Epi-AP, and equation 3 for A-AP and SAN-AP for $I_{\text{HERG}}$ peaks at APD$_{50}$ instead of APD$_{90}$ like Endo-AP, Epi-AP and Mid-AP, in these cells. Note that SDR2 was negative for depolarization.
- Using the values listed in Table 1, we calculated the $A$ values for Endo-AP, Mid-AP and Epi-AP (Fig. 7A), respectively, which are in agreement with the order of $I_{\text{HERG}}$ area (Fig. 7B). Interestingly, when the equation was applied to A-AP and SAN-AP, the resulting $A$ values also well reflect the relative size of $I_{\text{HERG}}$ area in these regions. This notion was further reinforced by the close correlation between the predicted and experimental $I_{\text{HERG}}$ areas (Fig. 7C).

Regional Differences of $I_{\text{HERG}}$ in Response to Dofetilide Blockade Revealed by AP-Clamp

Since the effect of $I_{Kr}$/I$\text{HERG}$ blockers is state-dependent with most of them generally acting on the open channels and since there is regional difference of APs in terms of the repolarization rate thereby the rate of HERG reopening, we reasoned that the effects of $I_{Kr}$/I$\text{HERG}$ blockers are different in cells from different regions of the heart. To test this notion, we went on to study the effects of dofetilide on $I_{\text{HERG}}$ using AP-clamp with AP waveforms collected from various regions. As illustrated in Fig. 9, 10 nM dofetilide which produced approximately 50±3% inhibition of $I_{\text{HERG}}$ area with Endo-AP reduced the current by 27±3%, 22±1%, 21±6%, and 30±3% with Mid-AP, Epi-AP, A-AP and SAN-AP, respectively (Fig. 9A and 9B). When the data were normalized to Epi-AP, it was clear that the degree of blockade of $I_{\text{HERG}}$ with A-AP was around 4-times greater than with Epi-AP and about 2–3-times greater than in APs of other regions (Fig. 9C). Also important is that there were transmural differences of $I_{\text{HERG}}$ in response to dofetilide block. For example, the blockade of $I_{\text{HERG}}$ with Endo-AP was 2.5-fold greater than with Epi-AP and 2-fold greater than with Mid-AP (Fig. 9C). Noticeably, these differential effects of dofetilide caused a remarkable change in the pattern of transmural difference of $I_{\text{HERG}}$; the normal $I_{\text{HERG}}$ gradient was changed to Mid-AP $>$ Epi-AP $>$ Endo-AP by dofetilide.
Figure 8. Influence of selective blockade of ion currents activating prior to \( I_{Kr} \) on \( i_{HERG} \) function reported by AP-clamp with atrial AP waveforms. A-D, A-AP configurations in the presence of selective blockers to inhibit \( I_{to} \), \( I_{Kur} \), or \( I_{CaL} \), from canine atrial myocytes, used to record the respective \( i_{HERG} \) waveforms. E, Mean data (\( n = 18 \) cells for each group) of area under \( i_{HERG} \) traces, \( I_{to} \) blocked by 500 nM nicotine; \( I_{Kur} \) blocked by 500 nM 4-AP; \( I_{CaL} \) blocked by 10 \( \mu \)M nisodipine. *\( p < 0.05 \) vs. Ctl. doi:10.1371/journal.pone.0072181.g008

Discussion

Novel Properties of HERG Channel Reported by AP-Clamp

The present study with AP-clamp revealed several important features of HERG \( K^+ \) channel function, which are otherwise masked with the conventional voltage pulse protocols. Some of these features have been described by previous work from other laboratories using AP-clamp techniques and therefore they will not be repeated in Discussion. Below are the selected aspects which we believe to represent the novel findings on HERG function revealed by the present study.

1. With AP-clamp, \( I_{HERG} \) in ventricular cells peaks at around \(-30 \) mV (\( V_{Peak} \)) and in atrial and SAN cells at \(-40 \) mV, with respect to voltage, as revealed by the instantaneous I–V relationships (Fig. 3). By comparison, \( I_{HERG} \) peaked at \(-10 \) mV with conventional pulse protocol. This implies that pulse protocols introduced \( \sim 20–30 \) mV voltage error to measurement of the voltage-dependence of HERG channel. The fully activated I–V relationship, obtained by various reactivating (repolarizing) steps following an inactivating (depolarizing) pulse with conventional pulse protocols or by ramp protocols, demonstrated the maximum \( I_{HERG} \) at \(-30 \) mV [21,23], approximating the AP-clamp value. Unfortunately, the protocol could not distinguish the regional difference of \( V_{Peak} \) or the voltage-dependence of HERG channel, especially the difference between V-APs and A-AP.

2. AP-clamp revealed that with respect to APD, \( I_{HERG} \) in ventricular cells peaked near \( APD_{50} \) and in atrial and SAN cells it peaked near \( APD_{70} \) (\( APD_{Peak} \)). The data imply that \( APD_{Peak} \) is predictable regardless of the length of APD. For example, the length of Mid-AP (\( APD_{50} \)) was around 400 ms and that of Endo-AP and Epi-AP was approximately 250 ms, but \( I_{HERG} \) in these regions all peaks at \( APD_{50} \) that corresponded to the length of phase 2 plateau. This was consistent with the view that \( I_{Kr} \) serves to terminate the phase 2 plateau of APs. Conventional pulse protocols do not provide

Table 3. Characteristics of the Atrial APs Used for AP-Clamp and the Corresponding \( I_{HERG} \) in the Presence of Varying \( K^+ \) Channel Blockers to Inhibit \( I_{to} \) or \( I_{CaL} \).

|         | Ctl | \( I_{to} \) | \( I_{Kur,d} \) | \( I_{CaL} \) |
|---------|-----|-------------|--------------|-------------|
| \( APD_{10} \) (mV–ms) | 27–17 | 13–34 | 26–25 | 25–15 |
| \( APD_{20} \) (mV–ms) | 16–30 | 3–60 | 15–51 | 14–21 |
| \( APD_{50} \) (mV–ms) | \( -19–82 \) | \( -28–107 \) | \( -20–86 \) | \( -20–46 \) |
| \( APD_{70} \) (mV–ms) | \( -30–96 \) | \( -38–122 \) | \( -32–93 \) | \( -31–56 \) |
| \( V_{Peak} \) (mV) | \( -41 \pm 1 \) | \( -43 \pm 1 \) | \( -49 \pm 1 \) | \( -40 \pm 1 \) |
| \( APD_{Peak} \) (% Repolarization) | \( -70 \pm 1 \) | \( -65 \pm 1 \) | \( -74 \pm 1 \) | \( -69 \pm 1 \) |
| \( A \) (pA/pF*\( s \times 1000 \)) | 11.1±0.6 | 9.6±0.5 | 13.4±0.9 | 8.2±0.7 |

\( V_{Peak} \): voltage at which \( V_{Peak} \) peaks; \( APD_{Peak} \) repolarization at which \( APD_{Peak} \) peaks; hence, the value 62% in the table can be expressed as \( APD_{Peak} \). A area under \( I_{HERG} \) trace, a measure of the volume of \( K^+ \) passing through HERG channel during an AP; \( V_{Peak} \): Voltage for half maximum inactivation calculated from the Boltzmann fit to the instantaneous inactivation curve; \( k \): slope factor from the Boltzmann fit for instantaneous conductance curve.

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Figure 9. Pharmacological property of \( i_{HERG} \) reported by AP-clamp. A, effects of \( I_{to} \)/HERG blocker dofetilide (Dof, 10 nM) on \( i_{HERG} \) with AP waveforms of various regions. *\( p < 0.05 \) vs. Ctl (\( n = 18 \) cells). B, % block of \( i_{HERG} \) over control by dofetilide with AP waveforms of various regions. *\( p < 0.05 \) vs. Endo-AP (\( n = 18 \) cells). C, Fold difference of \( i_{HERG} \) blockade by dofetilide with AP waveforms of various regions. Data were normalized to the value with Epi-AP.

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The initial rising phase (phase 1) of I_{HERG} was elicited by the depolarization phase of an AP and the major rising phases of I_{HERG} (phase 2 and phase 3) corresponded to the repolarization phases of an AP. The rate of rising phase was determined by the rate of membrane repolarization. No such conclusion can be drawn with conventional pulse protocols.

There was transmural heterogeneity of I_{HERG} function as indicated by the differences in I_{HERG} area, a measure of total K+ passing through HERG channels during an AP. The size of I_{HERG} area was found to be in the order of Endo-AP>Mid-AP>Epi-AP. Conventional pulse protocols would mask this important property of HERG channel.

There was regional difference of I_{HERG} in response to dofetilide blockade. Our results demonstrated that the effects of dofetilide on I_{HERG} elicited with APs from different regions of the heart were quantitatively different. The % blockade of I_{HERG} with A-AP was some 4-fold greater than Epi-AP and 2–3 fold greater than APs of other regions. The differences also existed among different layers of ventricular wall with significantly greater inhibition seen with Endo-AP than with Mid-AP and Epi-AP. Importantly, this transmural difference of I_{HERG} blockade by dofetilide broke the normal pattern of transmural gradient of I_{HERG} from Endo-AP>Mid-AP>Epi-AP under control conditions to Mid-AP>Epi-AP=Endo-AP in the presence of dofetilide. And the maximum transmural difference was also diminished from 6 (Endo-AP minus Epi-AP) to 2.5 (Mid-AP minus Epi-AP). Obviously, these differences cannot be revealed by using conventional pulse protocols.

During an AP, the complex nature of the interactions between different currents means that small changes in one current could alter the behavior of others. Indeed, our data show that I_{HERG} function was influenced by alterations of preceding currents. Our data show that I_{HERG} function varied in response to selective inhibition of any given current activated prior to I_{Ks}. For instance, blockade of I_{cal} markedly enhanced I_{HERG} and in contrast, blockade of I_{cal} caused pronounced reduction of I_{HERG} (Fig. 8).

### Biophysical Mechanisms for HERG Channel Reported by AP-Clamp

Evidently, HERG function is determined by HERG properties and AP characteristics; the latter indicates the differences in the rate of repolarization that governs the gating of HERG channels.

1. The dynamic changes of I_{HERG} (or the rising phase) during an AP were determined by rapid reactivation or removal of inactivation of HERG channels, a process depending upon the dynamic changes in rate of repolarization.

2. Voltage-dependent rapid reactivation of I_{HERG} determined the regenerative (all-or-none) repolarization. Under normal conditions, the cardiac repolarization process, once initiated, will carry on itself until it reaches full repolarization. The present study provides an explanation for the regenerative repolarization process: Membrane repolarization initiated by I_{o} activation and I_{cal} inactivation tends to reactivate I_{HERG}, and I_{HERG} reactivation in turn further repolarizes membrane which results in more I_{HERG} reactivation and further more repolarization.

3. The decaying phase of I_{HERG} with AP-clamp has been thought to be the balance between reactivation and deactivation of HERG channels [23,30]. Our data, however, provide an alternative mechanism for that. According to our data in Figure 2 as well as previous studies, no apparent deactivation process was seen at potentials positive to −50 mV and slow deactivation was noted at −60 mV to −80 mV with time constants of over 1 s. Considering that the decaying phase of I_{HERG} during a ventricular AP started from −40 mV and spanned <30 ms in duration, it is unlikely that deactivation process was involved. Similarly, changes in HERG conductance did not account for the decline either because during the whole decaying phase of I_{HERG} HERG conductance oppositely kept increasing (Fig. 6). In sharp contrast, the fall of driving force paralleled the declining of I_{HERG}. Thus, driving force or the electrochemical gradient may be the major factor determining I_{HERG} decay after the peak. This notion let us speculate that the rapid fall in driving force terminated repolarization and prevented excessive cardiac hyperpolarization that could otherwise happen with regenerative HERG reactivation during repolarization.

### Potential Implications of the Results

One of the important implications of our findings is that the repolarization-dependent reactivation of HERG channels contributing to the regenerative repolarization process of cardiac cells may help to suppress arrhythmias initiated by early afterdepolarizations and premature beats [11,23]. I_{HERG} reactivates to the maximum extent at around APD_{90}APD_{0} with respect to APD (APD_{peak}) and at −30 mV to −40 mV with respect to transmembrane potential (V_{peak}), which on one hand can avoid interfering with normal Ca\(^{2+}\) entry into the cells to initiate excitation-contraction coupling because I_{cal}, normally peaks at around −10 mV, and on the other hand can prevent excessive or secondary I_{cal} activation by generating large volume of outward K\(^{+}\) flow. This mechanism can serve to counteract any secondary depolarizations falling during this phase, particularly those generated by I_{cal}.

Another important implication of our findings is that there exist regional heterogeneity and transmural gradient of I_{HERG}, when assessed with I_{HERG} area. Given that myocytes from different layers of ventricle or from different regions of the heart bear the same I_{Ks} current densities [26], the differences of I_{HERG} area may
reflect the true regional heterogeneity and transmural gradient. These regional differences of \( I_{\text{HERG}} \) may partially account for the regional variations of cardiac repolarization and may have some important pathophysiological implications: being antiarrhythmic under physiological conditions but also being proarrhythmic or arrhythmogenic when in presence of HERG blockers which create transmural dispersion of repolarization. One example of the latter has been documented by Di Diego et al. [31] who showed that cisapride (a potent blocker of HERG channel) induced transmural dispersion of repolarization and *torsade de pointes* in the canine left ventricular wedge preparation during epicardial stimulation.

Also significant is that the extents of drug blockade of \( I_{\text{HERG}} \), specifically, the effects of dofetilide on \( I_{\text{HERG}} \) are different with APs from different regions. Dofetilide is currently in clinical use for suppressing atrial arrhythmias. Moreover, in the heart, the contribution of HERG to cardiac repolarization in any given cell type will also depend on the level of channel expression in that cell. A study reported by Gintant [32] suggests that in canine heart \( I_{Kr} \) density in atrium is approximately 50% higher than in ventricle. In terms of the transmural differences of dofetilide blockade of \( I_{\text{HERG}} \), our results show that dofetilide diminishes the maximum transmural differences of \( I_{\text{HERG}} \) area to 2.5 (Mid-AP minus Epi-AP) from 6 (Endo-AP minus Epi-AP) for control. Unfortunately, we are unable to draw any conclusions from the present study as to whether the reduction of the transmural difference of \( I_{\text{HERG}} \) will increase or decrease transmural dispersion of repolarization.

Finally, our data revealed for the first time the interactions between \( I_{Kr}/I_{\text{HERG}} \) and \( I_{to}, I_{Kur} \) or \( I_{CaL} \) that activates before \( I_{Kr}/I_{\text{HERG}} \) does. The important message from these results can be understood as a gate-safeguard mechanism of HERG function for cardiac repolarization. It is known that \( I_{Kur} \) activates rapidly upon membrane depolarization with a linear or slightly outward rectifying I–V relationship and stays activated with only partial inactivation during the whole course of an AP [25,29], indicating that it contributes to various phases of repolarization, particularly the phase 2 plateau. The present study demonstrates that when \( I_{Kur} \) is suppressed, \( I_{Kr}/I_{\text{HERG}} \) tends to increase. This can serve as a compensatory mechanism or a “repolarization reserve” to minimize the excessive slowing of repolarization due to \( I_{Kur} \) blockade. In sharp contrast, when \( I_{CaL} \) is depressed, \( I_{Kr}/I_{\text{HERG}} \) tends to decrease. This on one hand can allow a longer time for more \( Ca^{2+} \) entry for contractile function and on the other hand can limit the excessive acceleration of repolarization consequent to \( I_{CaL} \) reduction. Interestingly, \( I_{to} \) which contributes mainly to the initial repolarization, when blocked does not significantly alter \( I_{Kr}/I_{\text{HERG}} \) that has relatively small contribution to phase 1 repolarization. It is tempting to speculate that in atrial fibrillation when \( Ca^{2+} \) channel function is depressed [33], \( I_{Kur} \) will decrease accordingly so as to promote \( Ca^{2+} \) entry and minimize APD shortening.

**Limitations of the Study**

The assessment of \( I_{\text{HERG}} \) in the present study was based upon a few selected AP waveforms, which are representative of APs from different regions of the heart. However, it should be noted that the results may not be extrapolated to cells from other regions or subregions as AP waveforms are different in every different cells. Moreover, in native cells with co-existing multiple ion currents that can influence the function of HERG channels, the properties of \( I_{Kr} \) may not be the same as \( I_{\text{HERG}} \) recorded in a HERG-overexpressing cell line.

**Author Contributions**

Conceived and designed the experiments: ZPS. Performed the experiments: PZ, PG. Analyzed the data: PZ, PG, XLB. Contributed reagents/materials/analysis tools: XLB. Wrote the paper: ZPS.

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