Differences Between Ion Binding to eag and HERG Voltage Sensors Contribute to Differential Regulation of Activation and Deactivation Gating

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ABBRVIATIONS
eag ether-à-go-go
HERG human ether-à-go-go related gene
LQTS long QT syndrome
MiRP1 MinK-related protein 1
PCR polymerase chain reaction

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HERG (human ether-à-go-go related gene) (KCNH2) encodes two alternatively spliced voltage-gated K+ channel subunits (HERG1a and HERG1b) that co-assemble with MiRP1 (MinK-Related Protein 1) (KCNE2), a protein with a single transmembrane segment, to form the cardiac IKr channel.1,2 IKr channels play an essential role in repolarization of the cardiac action potential due to their unique gating kinetics. During action potential depolarization, IKr channels open and then very rapidly inactivate by a C-type mechanism.3-7 Rapid inactivation limits the availability of IKr channels during the plateau phase of the action potential. As the membrane begins to repolarize, however, IKr channels rapidly recover from inactivation and reenter an open, conducting conformation. Channel closure (deactivation) from this state occurs slowly. As a result, IKr channels generate a resurgent K+ current that reinforces and accelerates repolarization of the cardiac action potential. These unique gating properties of cardiac IKr channels are an intrinsic feature of heterotetrameric channels composed of the HERG1a and HERG1b subunits, although coassembly with MiRP1 subunits may modify the kinetics and pharmacology of HERG channels to more closely resemble those of native cardiac channels.1,8

The essential role of HERG channels in the heart is underscored by the finding that mutations in HERG are responsible for Long QT Syndrome Type 2 (LQTS2).3,9,10 In LQTS, a defect in repolarization of the cardiac action potential results in a lengthened QT interval detected by electrocardiogram. Individuals with LQTS are at increased risk for ventricular arrhythmia and sudden death.11 Although congenital forms of LQTS are rare, acquired LQTS is a significant health problem and is typically due to drug interactions with HERG channels.12,13 In some cases, naturally occurring polymorphisms in HERG and MiRP1 may be responsible for increased drug sensitivity.14,15

The process and kinetics of deactivation in HERG channels are central to the physiological role of IKr channels. Interestingly, the rate of deactivation in HERG channels is modulated by relatively small changes in extracellular Mg2+ or Ca2+ in their physiological concentration ranges. These ions appear to act by modifying channel gating rather than by blocking the pore or changing its conduction properties.16-19 We have tested the hypothesis that extracellular divalent cations regulate gating kinetics by binding directly to the voltage sensor in HERG channels. This hypothesis is predicated on our previous work.
with the ether-à-go-go (eag) K⁺ channel. In eag, extracellular Mg²⁺ regulates the kinetics of channel activation by binding to two acidic residues located in transmembrane segments S2 and S3.²⁰ Among voltage-gated K⁺ channels, these positions are occupied by acidic residues only in the eag subfamily, which explains why this form of ion regulation is uniquely found in this group. We now report that the HERG voltage sensor contains an ion-binding site that modulates gating kinetics. In HERG, ion binding affects the rates of both activation and deactivation, in contrast to eag, where effects on deactivation are not prominent.²¹,²² In HERG, the bound ion interacts with both eag specific and widely conserved acidic residues. This is in contrast to eag, where only eag subfamily-specific residues are implicated in ion binding.²⁰ We propose that differences between the ion-binding sites explain why ion binding prominently modulates deactivation gating in HERG but not in eag. Despite these differences, the data indicate that the presence of an ion-binding site between S2 and S3 is a conserved feature of members of the eag subfamily of voltage-gated K⁺ channels.

**EXPERIMENTAL PROCEDURES**

**Molecular biology.** The wild-type HERG1α cDNA clone was kindly provided by Dr. Mark Keating (Harvard Medical School, Boston, MA). Mutations in HERG1α were generated by the QuikChange method (Stratagene, La Jolla, CA) or by polymerase chain reaction (PCR) using a four primer strategy.²³ Mutated constructs were verified by Big Dye terminator sequencing (UCLA Sequencing Core Facility, Los Angeles, CA). Constructs in the pSP64 vector were linearized with EcoRI before in vitro transcription using the mMessage mMachine kit (Ambion, Austin, TX). RNA was injected into Xenopus oocytes for electrophysiological analysis.²⁴

**Electrophysiology.** Ionic currents from wild-type and mutant HERG channels were recorded at room temperature (20–22°C) 2–4 days after RNA injection using a two electrode voltage clamp.²⁴,²⁵ Electrodes were filled with 3 M KCl and had resistances of 0.3–1.0 MΩ. Oocytes were bathed in a modified ND96 solution containing 96 mM NaCl, 2 mM KCl, 5 mM HEPES, pH 7.5. The concentrations of MgCl₂ and CaCl₂ were varied as indicated in each experiment. The bath solution was changed using a gravity-driven perfusion system that was governed by solenoid valves and controlled manually.

Voltage pulse protocols were generated and data were acquired using pClamp software and a TL1 Labmaster interface (Axon Instruments, Foster City, CA). Data were sampled at 2–40 kHz and filtered at one fifth the sampling frequency using an 8-pole Bessel filter (Frequency Devices, Haverhill, MA). Linear capacitative and leak currents were subtracted using a P/4 protocol.²⁶

HERG ionic currents were evoked by pulsed from a holding potential of -80 mV to voltages ranging from -100 mV to +80 mV or to +100 mV in 20 mV increments. Tail currents were recorded upon repolarizing to -70 mV. The sampling rate was varied as needed to resolve deactivation kinetics. Tail currents were fitted with double or single exponential functions to derive time constants for deactivation (τ_deact). Activation kinetics was characterized by an envelope of tails procedure.²⁷ Values of τ_deact and τ_acq are provided as mean ± SEM. Statistical significance was assessed using a one-way ANOVA.

The probability of opening as a function of voltage was determined by measuring the amplitudes of isochronal tail currents evoked by repolarizing to -70 mV from various depolarized potentials. Tail current amplitudes were normalized to the maximum value and plotted versus pulse potential. The data were fitted with a single Boltzmann equation using Origin 7.0 (Origin Lab, Northampton, MA) to obtain slope and midpoint potential values.

**RESULTS**

Extracellular Mg²⁺ accelerates deactivation in wild-type HERG channels. HERG ionic currents were recorded in Xenopus oocytes in the presence or absence of 10 mM extracellular Mg²⁺ using a two-electrode voltage clamp (Fig. 1A). Currents were elicited by pulsing from a holding potential of -80 mV to voltages ranging from -100 to +80 mV. The membrane was then repolarized to -70 mV to record tail currents. Current amplitudes during the depolarizing pulses were low, as expected for the rapidly inactivating HERG channel.⁶ Upon repolarization and recovery from inactivation, however, large outward tail currents were recorded as the channel deactivated.

Extracellular Mg²⁺ significantly accelerated tail currents (Fig. 1A). The kinetics of deactivation at -70 mV were quantified by fitting two exponential components to the tail current records, providing values for two time constants, τ_fast and τ_slow (Table 1). In the presence of 10 mM MgCl₂, the values of τ_fast and τ_slow at -70 mV were reduced by approximately 2-fold and 3-fold, respectively (Table 1). Mg²⁺ also accelerated deactivation about 2-fold when HERG was co-expressed with MIRP1 (data not shown).

Extracellular Mg²⁺ slows activation in wild-type HERG channels. In eag, Mg²⁺ slows channel opening with little effect on channel closing.²¹,²²,²⁸,²⁹ In HERG channels, any effect of Mg²⁺ on activation kinetics is masked by the fast rate of inactivation. To investigate whether ion binding affects activation kinetics in HERG channels, we used an envelope of tails procedure (Fig. 1B).²⁷ To activate the channel, the membrane was stepped from -80 mV to +40 mV for pulses of increasing duration in the presence or absence of Mg²⁺. Peak tail current amplitudes were measured upon repolarization to -80 mV. The time course of ionic current activation was then inferred from the increase in tail current amplitude as a function of pulse duration (Fig. 1B). At +40 mV, 10 mM MgCl₂ slowed activation by approximately 2-fold (Table 2).

Our results indicate that extracellular Mg²⁺ modulates both activation and deactivation gating in HERG channels. Together, these effects reduce the maximum conductance measured in the presence of Mg²⁺ by about 20% (Fig. 1C).

Physiological concentrations of Mg²⁺ modulate HERG gating. To investigate the concentration dependence of Mg²⁺ modulation in HERG, deactivation kinetics were compared at Mg²⁺ concentrations up to 20 mM (Fig. 1D, left). Concentrations as low as 0.5 mM significantly accelerated deactivation; 10 mM Mg²⁺ gave a near maximal effect. To estimate the apparent affinity of HERG for Mg²⁺, changes in τ_fast and τ_slow in the absence and presence of Mg²⁺ were normalized (Δτ_fast/Δτ_max and Δτ_slow/Δτ_max) and plotted versus Mg²⁺ concentration (Fig. 1D, right). The data were fitted with the equation Δτ/Δτ_max = [Mg²⁺]/(K_m + [Mg²⁺]), providing estimates for the half maximal effective concentrations for Δτ_fast and Δτ_slow of 2.2 and 1.8 mM respectively. Given experimental variability, these values do not differ significantly (Fig. 1D). The steepest change in deactivation kinetics occurred between 0 and 3 mM. Since the
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Normal concentration of Mg\(^{2+}\) in serum ranges from 0.8 to 1.2 mM, Mg\(^{2+}\) modulation of deactivation in HERG channels is likely to be important physiologically.

**Extracellular Ca\(^{2+}\) modulates HERG gating.** In eag, the ion-binding site between S2 and S3 can accommodate different divalent cations, including Mg\(^{2+}\), Mn\(^{2+}\) and Ni\(^{2+}\), all of which modulate activation gating.\(^{20,30,31}\) To investigate whether multiple ions modulate HERG gating, we measured activation and deactivation kinetics in the presence and nominal absence of extracellular Ca\(^{2+}\) (Fig. 2). Like Mg\(^{2+}\), Ca\(^{2+}\) accelerated deactivation and slowed activation (Fig. 2).

Mg\(^{2+}\) binds between S2 and S3 to modulate deactivation and activation kinetics. The HERG protein contains a number of acidic residues in transmembrane segments S2 and S3 (Fig. 3A). Three of these (D456 and D466 in S2, D501 in S3) correspond to acidic residues that are widely conserved among members of the voltage-gated ion channel superfamily.\(^{32}\) Two others (D460 in S2 and D509 in S3) correspond to acidic residues that are conserved only among members of the eag subfamily of voltage-gated K\(^{+}\) channels (Fig. 3A).\(^{33,34}\) In eag, these subfamily-specific acidic residues contribute to a divalent ion-binding site that modulates activation gating.\(^{20,30,31,35}\)

To test the hypothesis that acidic residues in S2 and/or S3 contribute to an ion-binding site that modulates HERG gating, the five aspartate residues in S2 and S3 were mutated singly and in pairs to alanine. D501A in S3 eliminated functional expression and was not studied further. Other mutant constructs produced functional channels in Xenopus oocytes. Gating properties of the mutant channels were qualitatively similar to those of wild-type HERG, with small currents during depolarizing pulses and large tail currents evoked upon repolarization (data not shown). However, all of the mutant constructs deactivated more rapidly than wild-type HERG (Table 1). To determine whether the mutations reduced the effect

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**Figure 1.** Extracellular Mg\(^{2+}\) accelerates deactivation of wild-type HERG channels. (A) Wild-type HERG currents were recorded in the absence (left) or presence (middle) of 10 mM extracellular Mg\(^{2+}\). From a holding potential of -80 mV, 2 s test pulses ranging from -100 mV to +80 mV were applied in 20 mV increments prior to repolarization to -70 mV for 8 sec. Representative current traces are shown. Right, Tail currents evoked upon return from +60 mV to -70 mV in the absence [solid trace] or presence [dotted trace] of 10 mM Mg\(^{2+}\) have been scaled and superimposed to compare the time course of deactivation. Tail currents were fitted with two exponential components [dashed lines]. The bath solution contained 96 mM NaCl, 2 mM KCl, 0.5 mM CaCl\(_2\), 5 mM HEPES, pH 7.5. The pulse protocol is shown below the current traces. (B) Activation kinetics of wild-type HERG channels were measured using an envelope of tails protocol. Upper, Representative current traces were evoked at +40 mV by test pulses of different durations. Tail currents were recorded upon repolarization to -80 mV. Arrows indicate the peak tail currents. The pulse protocol is shown below the current traces. Lower, Normalized peak tail current amplitudes obtained in the presence [■] or absence [●] of 10 mM Mg\(^{2+}\) have been plotted versus pulse duration. Data are shown as mean ± SEM, n = 5. Each data set was fitted with a single exponential component to obtain values for \(\tau_{\text{fast}}\) in the presence (dashed curve) or absence (solid curve) of Mg\(^{2+}\). (C) \(I_{\text{tail,0}}/I_{\text{tail,max,0}}\) Mg [■] and \(I_{\text{tail,10}}/I_{\text{tail,max,10}}\) Mg [●] are plotted versus V. (D) Left, Values of the deactivation time constants \(\tau_{\text{fast}}\) [■] and \(\tau_{\text{slow}}\) [●] obtained upon return from +40 mV to -70 mV have been plotted versus Mg\(^{2+}\) concentration. The horizontal bar indicates the physiological range of Mg\(^{2+}\) concentration. Data are shown as mean ± SEM, n = 7. In this and subsequent figures, if error bars are not visible, they are smaller than the size of the symbol. Right, The difference between the fitted \(\tau_{\text{fast}}\) or \(\tau_{\text{slow}}\) values (\(\Delta\tau_{\text{fast}}\) [■] and \(\Delta\tau_{\text{slow}}\) [●]) obtained in the absence and presence of Mg\(^{2+}\) were normalized and plotted versus Mg\(^{2+}\) concentration. In this figure, \(\Delta\tau_{\text{fast}} = \tau_{\text{fast,0 Mg}} - \tau_{\text{fast,10 Mg}}\) and \(\Delta\tau_{\text{slow}} = \tau_{\text{slow,0 Mg}} - \tau_{\text{slow,10 Mg}}\). The half maximal effective concentration for Mg\(^{2+}\) was estimated as described in the text.
The presence or absence of 10 mM Mg\(^{2+}\) and tail currents were dramatically reduced in HERG wild-type channels was evident in the deactivation kinetics of several of the mutant constructs (Fig. 3B). Unlike wild-type HERG, some of the mutant channels had deactivation kinetics that were well-fitted by a single exponential component. Tail currents were fitted with one or two exponential components, as needed, to quantify deactivation kinetics in the presence and absence of Mg\(^{2+}\) (Fig. 4 and Table 1). In the single mutant constructs, the sensitivity of deactivation kinetics to Mg\(^{2+}\) was reduced to varying degrees, but the residual modulation was statistically significant (Table 1). For D456A, however, the effect of Mg\(^{2+}\) was significant only for the faster of two components of deactivation. Combination of D456A with D460A or D509A abolished Mg\(^{2+}\) modulation of deactivation kinetics. In contrast, D460A + D509A and D466A + D509A retained a small but statistically significant sensitivity to Mg\(^{2+}\). It is noteworthy that neutralization of both eag subfamily-specific acidic residues, D460 in S2 and D509 in S3, was not sufficient to eliminate Mg\(^{2+}\) modulation of deactivation kinetics. Rather, neutralization of either eag subfamily-specific acidic residue in combination with D456A was required. D456 in S2 corresponds to an acidic residue widely conserved among voltage-gated ion channels. In contrast, in eag, the analogous residue, D274, does not contribute to the ion-binding site.\(^{20,30}\) Our results are consistent with the conclusion that HERG channels contain an ion-binding site between S2 and S3 that modulates deactivation kinetics.

We also investigated whether mutating acidic residues in S2 and S3 affected Mg\(^{2+}\) modulation of activation kinetics (Fig. 5). Single and double mutant constructs, including D466A located near the cytoplasmic end of S2, dramatically reduced or abolished the slowing of activation kinetics mediated by extracellular Mg\(^{2+}\) (Fig. 5 and Table 2).

Mg\(^{2+}\) shifts Po-V curve in wild-type and mutant channels. Extracellular divalent ions have been reported to shift the voltage dependence of HERG activation in the depolarized direction.\(^{18,19,36,37}\) The size of the shift varies among different reports, perhaps because different ionic conditions were used.\(^{18,19,36,37}\) We found that extracellular Mg\(^{2+}\) shifted the voltage dependence of activation in the depolarized direction in wild-type channels (Fig. 6 and Table 3). To determine whether mutating acidic residues in S2 and S3 reduced or eliminated the effect of Mg\(^{2+}\) on the steady state properties of activation, we characterized the probability of opening as a function of voltage (Po-V) in mutant channels in the presence and absence of Mg\(^{2+}\). With the exception of D466A, the single and double mutations

### Table 1: Deactivation time constants for HERG and mutant constructs in the absence and presence of 10 mM Mg\(^{2+}\)

|                  | 0 mM Mg\(^{2+}\) | 10 mM Mg\(^{2+}\) | n  | V\(_{\text{test}}\) |
|------------------|------------------|------------------|----|------------------|
| WT (slow)**      | 2567 ± 129       | 789 ± 82         | 11 | +40 mV           |
| WT (fast)**      | 480 ± 25         | 219 ± 21         | 11 | +40 mV           |
| D456A (slow)     | 201 ± 21         | 165 ± 10         | 6  | +40 mV           |
| D456A (fast)**   | 52 ± 2           | 39 ± 1           | 6  | +40 mV           |
| D460A*           | 99 ± 5           | 76 ± 3           | 8  | +40 mV           |
| D466A*           | 111 ± 5          | 84 ± 7           | 5  | +40 mV           |
| D509A*           | 47 ± 1           | 42 ± 1           | 8  | +40 mV           |
| D456A/D460A (slow)| 163 ± 7          | 166 ± 9          | 8  | +60 mV           |
| D456A/D460A (fast)| 24 ± 1           | 26 ± 2           | 8  | +60 mV           |
| D456A/D509A      | 63 ± 3           | 59 ± 3           | 7  | +60 mV           |
| D460A/D509A*     | 50 ± 2           | 44 ± 1           | 5  | +60 mV           |
| D466A/D509A**    | 76 ± 2           | 53 ± 2           | 11 | +60 mV           |

Values for the deactivation time constants (in ms, mean ± SEM) were derived from single or double exponential fits to the tail currents (see Figs. 1A and 3B). The number of experiments, n, and the test pulse, V\(_{\text{test}}\), are indicated. * and ** denote significant differences in the presence and absence of 10 mM Mg\(^{2+}\) p < 0.001 and p < 0.05, respectively, using one-way ANOVA.

### Table 2: Activation time constants for HERG and mutant constructs in the absence and presence of 10 mM Mg\(^{2+}\)

|                  | 0 mM Mg\(^{2+}\) | 10 mM Mg\(^{2+}\) | n  | V\(_{\text{test}}\) |
|------------------|------------------|------------------|----|------------------|
| WT**             | 56 ± 5           | 136 ± 7          | 5  | +40 mV           |
| D456A            | 137 ± 14         | 166 ± 27         | 5  | +40 mV           |
| D460A            | 63 ± 5           | 69 ± 5           | 5  | +60 mV           |
| D466A            | 72 ± 2           | 86 ± 7           | 5  | +40 mV           |
| D509A            | 51 ± 6           | 62 ± 3           | 4  | +60 mV           |
| D456A/D460A      | 330 ± 32         | 422 ± 41         | 5  | +70 mV           |
| D456A/D509A      | 129 ± 15         | 154 ± 13         | 3  | +70 mV           |
| D460A/D509A*     | 146 ± 8          | 192 ± 15         | 5  | +70 mV           |
| D466A/D509A**    | 194 ± 13         | 203 ± 16         | 6  | +70 mV           |

Values for the activation time constants (in ms, mean ± SEM) were derived from single exponential fits to the currents evoked by an envelope of tails protocol (see Figs. 1B and 5A). The number of experiments, n, and the test pulse, V\(_{\text{test}}\), are indicated. * and ** denote significant differences in the presence and absence of 10 mM Mg\(^{2+}\) p < 0.001 and p < 0.05, respectively, using one-way ANOVA.
shifted the voltage dependence of HERG activation to more depolarized potentials in the absence of Mg$^{2+}$. An additional positive shift was seen when 10 mM Mg$^{2+}$ was added to the bath solution (Fig. 6 and Table 3). Thus, the effect of Mg$^{2+}$ on the voltage dependence of activation was not primarily due to ion binding between S2 and S3. It is likely that Mg$^{2+}$ exerted a non-specific surface charge effect on the voltage sensor in both wild-type and mutant channels.8 In our experiments, the bath solution contained 0.5 mM Ca$^{2+}$ with or without 10 mM Mg$^{2+}$. Adding Ca$^{2+}$ improved the quality of our electrophysiological recordings and served to partially offset surface charge effects due to added Mg$^{2+}$. However, in all likelihood Ca$^{2+}$ binds to the HERG voltage sensor, accounting for its effects on opening and closing kinetics (see Fig. 2). The use of 0.5 mM Ca$^{2+}$ was therefore a compromise, reducing the effect of Ca$^{2+}$ on gating kinetics in Mg$^{2+}$-free recordings, while at the same time stabilizing the membrane and providing some surface charge screening. Not surprisingly, 0.5 mM Ca$^{2+}$ was insufficient to screen entirely the non-specific surface charge effects of 10 mM Mg$^{2+}$. We therefore focused on the kinetic effects of Mg$^{2+}$ to identify the ion-binding site in the HERG voltage sensor domain.

**DISCUSSION**

Modulation of HERG gating by extracellular divalent cations may help counteract detrimental cardiac effects of hypomagnesemia and hypocalcemia. Modulation of HERG gating by Mg$^{2+}$ and Ca$^{2+}$ is likely to be physiologically important because a major target of ion regulation is the slow deactivation process, which is a critical factor in repolarization of the cardiac action potential.39,40 Furthermore, significant changes in deactivation kinetics result from small changes in extracellular Mg$^{2+}$ and Ca$^{2+}$ in and near the normal concentration range. Hypomagnesemia and hypocalcemia increase the excitability of the heart and contribute to cardiac arrhythmia.41 However, such changes would also slow the rate of HERG deactivation, leading to additional outward K$^+$ current that would be expected to counteract the hyperexcitability. Therefore, ionic modulation of deactivation gating in HERG may serve as a partially protective mechanism in the heart.

Our experiments were conducted using the HERG1a splice variant, whereas cardiac channels contain HERG1a and 1b subunits.2 HERG1a and 1b are identical except at the amino terminus.8 Both subunits contain the ion-binding site residues in the voltage sensor, consistent with the idea that divalent ions would accelerate deactivation in native cardiac channels. It is worth noting that the deactivation kinetics of heterotetrameric HERG1a/HERG1b and homotetrameric HERG1a channels differ.9 Therefore deactivation kinetics in HERG1a/HERG1b channels measured in the presence and absence of extracellular divalent ions would be expected to deviate from the values reported in this study.8

Mg$^{2+}$ interacts with D456, D460 and D509 to modulate deactivation gating in HERG. The prominent modulation of activation and deactivation in HERG channels indicates that extracellular divalent cations are able to bind to voltage sensors in the resting conformation to affect the rate of opening and in the activated conformation to affect the rate of closing. Our data support the conclusion that Mg$^{2+}$ interacts with D456 and D460 in S2 and D509 in S3 to accelerate channel closing. This conclusion is compatible with the results of Fernandez et al.57 They found that the non-physiological ion Cd$^{2+}$ binds to a site formed by the same three residues and thereby shifts the voltage-dependence of activation. Fernandez and co-workers measured shifts in activation caused by 0.5 mM Cd$^{2+}$ in the constant presence of 1 mM Ca$^{2+}$ and 2 mM Mg$^{2+}$. The presence of excess Ca$^{2+}$ and Mg$^{2+}$ presumably reduced non-specific surface charge effects caused by Cd$^{2+}$ binding to negatively charged lipid headgroups or acidic amino acids located in extracellular loops,38 making it feasible to measure shifts in activation caused by Cd$^{2+}$ binding to the voltage sensor site. In contrast, much of the Mg$^{2+}$-induced shift in the voltage dependence of activation measured in our experiments is likely to result from a non-specific surface charge effect rather than specific-binding to the HERG voltage sensor.

Mg$^{2+}$ binding to the resting conformation may depend on the negative electrostatic potential of the cleft between S2 and S3. Our data suggest that Mg$^{2+}$ also binds between S2 and S3 to slow channel opening. Interestingly, the mutations had differential effects on Mg$^{2+}$ modulation of activation and deactivation gating. Whereas only
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Results suggest that divalent ions affect HERG activation and deactivation kinetics by different mechanisms. On the basis of cysteine modification experiments, Tseng and colleagues have proposed that HERG contains a long, externally-facing aqueous cleft that extends to D466 in S2 in the resting conformation. Our results suggest that binding of divalent cations to the resting voltage sensor depends more on the negative electrostatic potential of this cleft than on interactions with specific acidic residues. Reduction of the negative potential by neutralization of even one acidic residue appears to be sufficient to eliminate ion binding to the resting conformation. In contrast, our work on eag did not reveal a similar mechanism in that channel, likely due to subtle structural differences between the clefts in eag and HERG.

Differential regulation of activation and deactivation by ion binding in eag and HERG. Divalent cation modulation of deactivation in HERG and activation in eag are mechanistically related because both phenomena involve ion binding to a specific site that includes the eag subfamily-specific acidic residues in S2 and S3. In HERG, the widely conserved acidic residue D456 is also involved in ion binding, whereas the analogous residue in eag is not. We propose that differences in the eag and HERG binding sites contribute to the differential regulation of opening and closing kinetics in these channels.

High resolution structural information is not available for eag or HERG. However, the structures of voltage-gated K+ channels are likely to be highly conserved. To compare the eag and HERG binding sites, we generated tentative models based on the X-ray structure of a chimeric K+ channel in which the voltage sensor paddle of Kv2.1 has been inserted into Kv1.2 (Fig. 7). Although the sequences of HERG and eag diverge from that of the chimeric channel, S2 and S3 can be aligned unambiguously due to the presence of highly conserved residues (Fig. 3A).

To model the binding sites, residues I230 in S2 and Y267 in S3 in the chimeric channel sequence were changed to aspartate residues (Figs. 3A and 7A). For the HERG model, the conserved S2 residue E226 was also changed to an aspartate residue (Figs. 3A and 7B). Using numbers corresponding to the chimeric sequence, the eag binding site comprises D230 and D267, whereas the HERG binding site comprises D226, D230 and D267 (Fig. 7A and B). For simplicity,
we assumed that Mg\(^{2+}\) binds equidistant from the \(\gamma\)-carbon atoms of the aspartate residues that contribute to the binding site.

In the eag model, Mg\(^{2+}\) is located 3.8 Å from the \(\gamma\)-carbons of D230 and D267 in excellent agreement with equivalent distances in divalent cation binding sites of known structure (Fig. 7A).\(^{43-46}\) Mg\(^{2+}\) binds closer to the extracellular side of the membrane in HERG than in eag due to the participation of D226 in the binding site (Fig. 7B). In the HERG model, the distance between a centrally-located Mg\(^{2+}\) ion and the \(\gamma\)-carbons of D226, D230 and D267 is 8.2 Å, placing the carbonyl oxygens too far away for optimal coordination. This suggests that the structures of HERG and the chimeric channel differ in the vicinity of the binding site or that ion binding in HERG draws the coordinating residues closer together.

Previous work suggests a mechanistic explanation for the differential modulation of gating in HERG and eag. In K\(^{+}\) channels, the first four arginine residues in S4, called R1-R4, carry the bulk of the gating charge across the transmembrane electric field during voltage-dependent gating.\(^{47-49}\) Based on data obtained with Shaker and eag channels, we proposed a model for voltage sensor conformational changes that occur late during the activation process.\(^{30,50}\)

In this model, R3 and R4 move sequentially into the extracellular cleft where they interact with the widely conserved acidic residue in S2 (Fig. 3A).\(^{30,50}\) This residue, corresponding to D456 in HERG, D274 in eag and E226 in the chimeric channel, will be referred to as D/E1 to simplify the following discussion. In our model, D/E1 in S2 and R4 in S4 interact at short range in the activated confirmation.\(^{30,50}\) This electrostatic interaction has since been confirmed by the X-ray structures of K1.2 and the chimeric channel in the activated state.\(^{42,51,52}\) Significantly, the D/E1-R4 interaction is stable in molecular dynamics simulations using the K1.2 structure embedded in a lipid bilayer.\(^{53}\)

### Table 3 Activation parameters for HERG and mutant constructs in the absence and presence of 10 mM Mg\(^{2+}\)

| Construct | \(\Delta V_\text{1/2}\) (mV) | \(n\) | Number of Experiments |
|-----------|-----------------|------|----------------------|
| WT        | -22 ± 3         | 5    | 12 mV               |
| D456A     | 7 ± 1           | 6    | 8 mV                |
| D460A     | -4 ± 1          | 6    | 16 mV               |
| D466A     | -29 ± 2         | 5    | 19 mV               |
| D509A     | 14 ± 1          | 6    | 13 mV               |
| D456A/D460A | 55 ± 2       | 6    | 4 mV                |
| D460A/D509A | 39 ± 2       | 5    | 5 mV                |
| D460A/D509A | 42 ± 1       | 5    | 8 mV                |
| D466A/D509A | 34 ± 1       | 6    | 7 mV                |

Isochronal tail current amplitudes in the presence or absence of Mg\(^{2+}\) were measured at -70 mV and normalized to the maximum tail current amplitude obtained in the absence of Mg\(^{2+}\). Values for the midpoint potential (\(V_\text{1/2}\)) (in mV, mean ± SEM) were derived from plots of \(I_{\text{tail}}/I_{\text{tail,max}}\) versus \(V\) fitted with a Boltzmann equation of the form: \(I_{\text{tail}}/I_{\text{tail,max}}=1/(1+\exp[\Delta V_{1/2}(V)/A])\). The number of experiments, \(n\), and the shift of \(V_{1/2}, \Delta V_{1/2}\) induced by the presence of 10 mM Mg\(^{2+}\) are provided.

### Figure 7. Speculative models for the ion-binding sites in eag and HERG.

Models for (A) eag and (B) HERG were based on the X-ray structure of a chimeric K\(^{+}\) channel in an activated conformation.\(^{42}\) S1 (blue, corresponding to residues P161-E183 in the chimeric channel), S2 (pale green, residues E226-D244), S3 (pink, residues N253-S275) and S4 (cyan, residues V288-S307) are shown in ribbon form. The side chains of I230 in S2 and Y267 in S3 have been changed to aspartate. Additionally, E226 has been changed to aspartate in the HERG model. A Mg\(^{2+}\) ion (green sphere) is shown equidistant from the \(\gamma\)-carbons of the acidic residues that contribute to each binding site. Accession number for the structure of the chimeric channel in the protein data base is 2R9R. Models were generated using Win-coot v.0.3.3 and PyMOL.

### Figure 6. Mg\(^{2+}\) shifts the voltage dependence of activation in both wild-type and mutant channels. Upper: Isochronal tail current amplitudes in the presence or absence of Mg\(^{2+}\) were measured at -70 mV and normalized to the maximum tail current amplitude obtained in the absence of Mg\(^{2+}\). \(I_{\text{tail}}/I_{\text{tail,max}}\) has been plotted versus voltage for wild type (■,■); D456A (●,●); D460A (▲,▲); D466A (▼,▼); and D509A (♦,♦) channels in the absence (filled symbols) or presence (open symbols) of 10 mM Mg\(^{2+}\). Bottom: \(I_{\text{tail}}/I_{\text{tail,max}}\) has been plotted versus voltage for wild type (■,■); D456A + D460A (●,●); D456A + D509A (▲,▲); D460A + D509A (▼,▼); and D466A + D509A (♦,♦) channels in the absence (filled symbols) or presence (open symbols) of 10 mM Mg\(^{2+}\). The bath solution contained 96 mM NaCl, 2 mM KCl, 0.5 mM CaCl\(_2\), 5 mM HEPES, pH 7.5. After repolarization from various test potentials to -70 mV, isochronal tail current amplitudes were measured and normalized to the maximum amplitude obtained in the experiment. Data points represent the average of 5–6 experiments. Error bars are omitted for clarity. Each data set was fitted with a single Boltzmann function (solid curves, 0 mM Mg\(^{2+}\); dotted curves, 10 mM Mg\(^{2+}\)) to obtain values for the midpoint potential (\(V_{1/2}\)) and slope factor (see Table 3 legend). \(V_{1/2}\) values are provided in Table 3.
Ion binding in eag slows the rate determining step in activation, during which R3 moves into proximity with D/E1 to generate an intermediate closed state. In eag, D/E1 does not contribute to the binding site, indicating that bound ions are located below the position of D/E1 (see Fig. 7). We propose that bound ions reduce the negative electrostatic potential near the bottom of the cleft, slowing the rate determining transition for activation in which R3 enters the cleft to move towards D/E1.

In contrast, ion binding in HERG accelerates deactivation. Channel closing involves S4 movement, consistent with the observation that ionic tail currents and off gating currents have very similar kinetics in a variety of voltage-gated channels, including Shaker and eag. As the channel closes, S4 moves inward, disrupting the electrostatic interaction between D/E1 and R4. Because D/E1 contributes to the binding site, the electrostatic interaction between R4 and D/E1 would be weakened in the presence of a divalent ion, destabilizing the activated conformation and accelerating deactivation.

In conclusion, we propose that the location of the bound ion in the cleft plays a major role in determining whether divalent cations affect primarily the rate-determining step for opening, as in eag, or the closing transition that blocks ion conduction through the pore, as in HERG. This difference between the channels is intriguing given the essential role of HERG deactivation in heart function.

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