Molecular basis of altered hERG1 channel gating induced by ginsenoside Rg3

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**Abbreviations:** $g/g_{\text{max}}$, relative conductance; hERG1, human ether-à-go-go-related gene type 1; $I_{\text{peak}}$, peak outward current during test pulse; $I_{\text{tail}}$, tail current; $I_{\text{tail-peak}}$, peak tail current; $k$, slope factor of Boltzmann function; LQTS, long QT syndrome; MD, molecular dynamics; Rg3, 20(S)-ginsenoside Rg3, (2S,3R,4S,5S,6R)-2-[(2R,3R,4S,5S,6R)-4,5-dihydroxy-2-[[3S,5R,8R,9R,10R,12R,13R,14R,17S)-12-hydroxy-17-[(2S)-2-hydroxy-6-methylhept-5-en-2-yl]-4,4,8,10,14-pentamethyl-2,3,5,6,7,9,11,12,13,15,16,17-dodecahydro-1H-cyclopenta[alphenanthren-3-yl]oxy]-6-(hydroxymethyl)oxan-3-yl]oxy-6-(hydroxymethyl)oxane-3,4,5-triol; $V_h$, holding potential; RPR, RPR-260243, (3R,4R)-4-[3-(6-methoxy-quinolin-4-yl)-3-oxo-propyl]-1-[3-(2,3,5 trifluorophenyl)-prop-2-ynyl]piperidine-3-carboxylic acid; $V_{\text{ret}}$, return potential; VSD, voltage sensor domain; $V_t$, test potential; $V_{0.5}$, half-point (in mV) of Boltzmann function; WT, wild-type; $\tau_{\text{deact}}$, time constant of current deactivation
ABSTRACT

Outward current conducted by human ether-à-go-go-related gene type 1 (hERG1) channels is a major determinant of action potential repolarization in the human ventricle. Ginsenoside 20(S)-Rg3 (Rg3), an alkaloid isolated from the root of *Panax ginseng* slows the rate of hERG1 deactivation, induces channels to open at more negative potentials than normal and increases current magnitude. The onset of Rg3 action is extremely fast, suggesting that it binds to an extracellular accessible site on the channel to alter its gating. Here we used a scanning mutagenesis approach to identify residues in the extracellular loops and transmembrane segments of hERG1 that might interact with Rg3. Single or multiple residues of hERG1 were mutated to Ala or Cys and the resulting mutant channels were heterologously expressed in *Xenopus* oocytes. The effects of Rg3 on the voltage dependence of activation and deactivation rate of mutant channel currents were characterized using two-microelectrode voltage clamp. Mutation to Ala of specific residues in the S1 (Tyr420), S2 (Leu452, Phe463) and S4 (Ile521, Lys525) segments partially inhibited the effects of Rg3 on hERG1. The double mutant Y420A/L452A nearly eliminated the effects of Rg3 on voltage dependent channel gating, but did not prevent the increase in current magnitude. These findings together with molecular modeling suggest that Rg3 alters the gating of hERG1 channels by interacting with and stabilizing the voltage sensor domain in an activated state.


**Introduction**

Repolarization of the cardiac action potential is mediated in part by voltage-dependent, K⁺-selective hERG1 channels. Pharmacological block of these channels can prolong the action potential duration (APD) in ventricular myocytes and increase the risk of cardiac arrhythmia, in essence mimicking the phenotype of congenital long QT syndrome (LQTS). This serious cardiac side effect was found to be caused by so many common medications that preclinical screening of compounds for hERG1 channel activity was adopted as a routine assay during early stages of the drug discovery process. Extensive screening of compound libraries serendipitously resulted in the discovery of several compounds that activate rather than inhibit hERG1 channels (Gerlach, et al., 2010, Kang, et al., 2005, Zhou, et al., 2005). The hERG1 activators discovered to date alter channel gating by one or more molecular mechanisms, including an increase in single channel open probability, inhibition of inactivation, slowing of deactivation and shifting the voltage dependence of activation to more negative potentials (Sanguinetti, 2014).

In an attempt to relate mechanisms of action of hERG1 activators to channel structure, we have used a site-directed mutagenesis approach to identify residues that contribute to the putative binding site of these compounds. These studies suggest that RPR-260243 (Wu, et al., 2015), PD-118057 and ICA-105574 (Wu, et al., 2014) bind with four-fold symmetry to specific residues in the pore domain (S5-S6 region) to alter gating of the channel.

Rg3 is a large (785 Da) steroid glycoside isolated from the root of *Panax ginseng* that alters hERG1 gating by slowing the rate of channel closure (deactivation) and causing a leftward shift in the voltage dependence of channel activation (Choi, et al., 2011a, Wu, et al., 2016). The onset of these effects is extremely fast, suggesting that this large compound exerts its actions by binding to an extracellular accessible site. A previously published study reported that S631C
hERG1 channels were resistant to the effects of Rg3, leading to the suggestion that Ser631 is a major structural determinant of Rg3 interaction with the channel (Choi, et al., 2011a). Here, we reexamine the importance of Ser631 in mediating the effects of Rg3 on channel gating and search for other potential interacting residues using a thorough mutagenesis scan of residues in the hERG1 subunit that could be accessible by a large molecule from the extracellular space.

**Materials and Methods**

**Molecular Biology.** *HERG1* isoform 1a (*KCNH2*, NCBI Reference sequence: NM_000238) was cloned into pSP64 oocyte expression vector. Single or multiple point mutations in *KCNH2* were introduced into wild-type (WT) channels using the QuikChange site-directed mutagenesis kit (Agilent Technologies, Santa Clara, CA). To prepare cRNA for use in oocyte expression studies, pSP64 plasmids were linearized with EcoRI prior to *in vitro* transcription using the mMessage mMACHINE SP6 transcription kit (Thermo Fisher Scientific, Waltham, MA).

**Isolation and Voltage Clamp of Oocytes.** *Xenopus laevis* oocytes were harvested as previously described (Garg, et al., 2012) using methods approved by the University of Utah Institutional Animal Care and Use Committee. Isolated oocytes were incubated at 17°C in Barth's solution and studied 2 to 7 days after injection with 1-40 ng KCNH2 cRNA encoding WT or mutant hERG1 channels. Whole cell currents were measured by using standard two-electrode voltage-clamp techniques (Stühmer, 1992). Agarose cushion electrodes (Schreibmayer, et al., 1994) with a tip resistance of 0.5 to 1.5 MΩ when backfilled with 3 M KCl were used to minimize leakage of pipette solution into oocytes. Data acquisition was performed with a Dell personal computer, and Digidata 1322A data acquisition system, GeneClamp 500 amplifier controlled by pCLAMP 8.2 software (all Molecular Devices, Inc., Sunnyvale, CA).
For most experiments, hERG1 channel currents were activated with 4 s voltage steps to a test potential \( V_t \) that was varied from \(-80 \) to \(+40 \) mV in 10 mV increments from a holding potential \( V_h \) of \(-70, -80 \) or \(-90 \) mV. To compare current-voltage relationships before and after treatment of an oocyte with Rg3, the time-dependent outward current \( I_{\text{peak}} \) was measured at the end of each test pulse, normalized to the largest value determined under control conditions and plotted as a function of \( V_t \). Tail currents \( I_{\text{tail}} \) were measured immediately after a test pulse by stepping the membrane voltage to a return potential \( V_{\text{ret}} \) of \(-70 \) mV. The pulse to \(-70 \) was usually 4 s in duration but was extended to 10 s for slowly deactivating mutant channels. Before each test pulse a brief prepulse to \(-70 \) was applied and the magnitude of leak current at this potential was measured and subtracted from \( I_{\text{tail}} \) to quantify the peak tail current \( I_{\text{tail-peak}} \).

To estimate the fold-change in \( I_{\text{tail-peak}} \) induced by Rg3, a \( V_t \) of sufficient amplitude and duration (e.g., 1 or 4 s at \(+40 \) mV) to fully activate channels was applied, followed by a return pulse to \(-70 \) mV. Leak-subtracted \( I_{\text{tail}} \) at \(-70 \) mV was fitted with a single or double exponential function and extrapolated to the moment of membrane repolarization to estimate \( I_{\text{tail-peak}} \). This method was especially important for fast deactivating mutant channels where \( I_{\text{tail-peak}} \) could be severely underestimated due to insufficient time for channels to recover from inactivation. To measure the kinetics of \( I_{\text{tail}} \) over a wide voltage range, Y420A/L452A hERG1 channels were activated with a 1 s pulse to 40 mV followed by stepping to a \( V_{\text{ret}} \) that was varied in 10 mV increments from 20 to \(-130 \) mV.

**Solutions.** Barth's solution contained (in mM): 88 NaCl, 1 KCl, 0.41 CaCl\(_2\), 0.33 Ca(NO\(_3\))\(_2\), 1 MgSO\(_4\), 2.4 NaHCO\(_3\), 10 HEPES, 1 pyruvate, plus gentamycin (50 mg/liter), amikacin (100 mg/liter) and ciprofloxacin (25 mg/liter); pH adjusted to 7.4 with NaOH. For voltage clamp experiments, oocytes were bathed in an extracellular saline solution that contained (in mM): 96
NaCl, 2 KCl, 1 CaCl₂, 1 MgCl₂, 5 HEPES; pH adjusted to 7.6 with NaOH. Ginsenoside 20(S)-Rg3 (Sigma-Aldrich, St. Louis) was dissolved in DMSO to make a 1 or 10 mM stock solution that was stored at −20°C. Before each experiment, a stock solution was diluted with the extracellular saline solution to produce the desired final concentration of Rg3.

**Data Analysis.** PCLAMP 8.2 software (Molecular Devices, Sunnyvale, CA) was used to analyze the magnitude and kinetics of digitized currents. Origin 8.6 software (OriginLab Corp, Northampton, MA) was used to prepare graphs and figures. Deactivation of hERG1 channels was quantified by fitting the time-dependent decay of $I_{\text{tail}}$ to a mono- or bi-exponential function:

$$I_{\text{tail}}(t) = A_f e^{-t/\tau_f} + A_s e^{-t/\tau_s} + C$$

where $\tau_f$ and $\tau_s$ are the fast and slow time constants and $A_s/(A_s + A_f)$ is the relative amplitude of the slow component of deactivation. Deactivation of WT hERG1 channel $I_{\text{tail}}$ is a biexponential process, but when slowed by Rg3 a single exponential adequately describes the majority of current decay (Wu, et al., 2016). To simplify comparison of deactivation rates before and after treatment with Rg3, $I_{\text{tail}}$ at −70 mV was fitted with a single exponential function, and the fold-change in the deactivation time constant ($\tau_{\text{deact}}$) was calculated for each mutant channel type.

The relative conductance-voltage ($g/g_{\text{max}}-V_t$) relationship for hERG1 was determined by plotting normalized peak tail currents ($I_{\text{tail-peak}}$) versus $V_t$ for each cell. Origin 8.6 software was used for fitting the data with a Boltzmann function:

$$\frac{g}{g_{\text{max}}} = \frac{1}{1 + e^{(V_{0.5}-V_t)/k}}$$

where $V_{0.5}$ is the voltage required for half maximal $I_{\text{tail-peak}}$ activation and $k$ is the slope factor for the relationship. $V_{0.5}$ and $k$ values were determined for each individual cell and the averaged data for all the channels characterized in this study are presented in Supplemental Table 1. The fold-change in the maximum values of $I_{\text{tail-peak}}$ induced by Rg3 treatment for WT and mutant channels...
are presented in Supplemental Table 2. Data are expressed as mean ± S.D. \((n = \text{number of oocytes})\).

**Molecular Modeling.** A homology model of the voltage sensing domain (VSD) in the activated state was built using the recently solved cryo-EM structure of the rEAG1 channel (PDB 5K7L; Whicher and MacKinnon, 2016) as template and the Modeller9v14 program (Marti-Renom, et al., 2000). Since this model is not fully consistent with data on charge-pairings stabilizing the open state, an additional model (denoted activated\(_1\)), with the alignment shifted +3 was built, consistent with experimental data for hERG1 (Zhang, et al., 2005).

Coordinates of Rg3 were optimized with the Hartee-Fock 6-31G* basis set implemented in Gaussian09 (Frisch, et al., 2009). Docking was performed using the program Gold 4.0.1 (Cambridge DataCentre, Cambridge, UK) (Jones, et al., 1995). Coordinates of the geometric center calculated among residues Y420, L452, F463, I521 and K525 were taken as binding site origin. The binding site radius was set to 20 Å. 150,000 operations of the GOLD genetic algorithm were used to dock the compound into the two different VSD models. The best snapshots (highest binding score) were subsequently relaxed by molecular dynamics (MD) simulations.

MD simulations were performed in Gromacs5.1.4 and the amber99sb force field (Hornak, et al., 2006). VSD domains were embedded in an equilibrated simulation box consisting of 118 1-Palmitoyl-2-oleoylphosphatidylcholine (POPC) lipids and the SPC/E water model was used (Berendsen, et al., 1987). Electrostatic interactions were calculated explicitly at a distance <1 nm and long-range electrostatic interactions were calculated at every step by particle-mesh Ewald summation (Darden, et al., 1993). Lennard-Jones interactions were calculated with a cutoff of 1 nm. All bonds were constrained by using the LINCS (Linear Constraint Solver) algorithm (Hess,
et al., 1997), allowing for an integration time step of 2 femtoseconds. The Nose-Hoover thermostat (Nose, 1984) was used for temperature coupling ($\tau = 0.1$ picoseconds), and the Parrinello-Rahman barostat algorithm (Parrinello and Rahman, 1981) was used for pressure coupling. One thousand conjugate gradient energy-minimization steps were performed, followed by 5 nanoseconds of equilibration, in which the protein atoms were restrained with a force constant of $1000 \text{kJ} \times \text{mol}^{-1} \text{nm}^{-2}$ to their initial position, and ions, lipids, and solvent were allowed to move freely. Each system was subjected to 2 x 50 nanoseconds of unrestrained MD, during which coordinates were saved every 20 picoseconds for analysis.

**Results**

**The onset of Rg3 is rapid.** As reported previously (Choi, et al., 2011a, Wu, et al., 2016), the most obvious effect of Rg3 on hERG1 channels is a slowing in the rate of $I_{\text{tail}}$ deactivation and an altered voltage dependence of channel activation. Slowed deactivation is illustrated in Fig. 1A where WT hERG1 channels were activated by 4 s pulses to a range of test voltages and $I_{\text{tail}}$ was measured at a fixed $V_{\text{ret}}$ of $-70$ mV. Rg3 increased the magnitude of $I_{\text{peak}}$ at $V_t < 0$ mV, consistent with a negative shift in the voltage dependence of channel activation. This effect was confirmed in Fig. 1C where $I_{\text{tail-peak}}$ was plotted as a function of $V_t$. The $V_{0.5}$ for activation determined by fitting this relationship with a Boltzmann function was shifted from $-31.5$ mV under control conditions to $-42.6$ mV in the presence of 3 µM Rg3, a maximally effective concentration (Choi, et al., 2011a, Wu, et al., 2016). In addition, Rg3 increased the maximum amplitude of $I_{\text{tail-peak}}$ by $10 \pm 4.5\%$ ($n = 5$), an effect likely due to an increase in channel open probability because the alkaloid has no effect on the voltage dependence of inactivation (Wu, et al., 2016). The kinetics of Rg3-induced changes in channel gating were probed by applying repetitive 1 s voltage steps,
once every 5 s, to a $V_t$ of 0 mV from a $V_h$ of −90 mV before and after rapidly switching the perfusate from the control solution to a solution containing 3 µM Rg3 (Fig. 1D). After the addition of Rg3, current was increased midway through the first test pulse and $I_{\text{tail}}$ measured at −70 mV was dramatically slowed. $I_{\text{tail}}$ slowed further during the second pulse, but was not further changed by continued pulsing. The magnitude of $I_{\text{tail}}$ was measured at 1 s after stepping to −70 mV to estimate changes in deactivation kinetics in this and 6 other oocytes. The normalized $I_{\text{tail}}$ at 1 s was plotted as a function of time after application of Rg3 and fitted with a single exponential function (Fig. 1E). The calculated $\tau$ of 1.9 s is only an approximation given the solution flow rate of 2 mL/min and a chamber volume of 0.2 mL. We previously reported that the onset and recovery from modification of hELK1 channel gating by Rg3 (0.3 µM) was also very rapid with a $\tau$ of 6.1 s and 19.3 s, respectively (Wu, et al., 2016). By comparison, hERG1 blockers bind to the central cavity and therefore must cross the cell membrane to gain access to the central pore. The rates of block onset by compounds that must cross the oocyte membrane to reach their putative receptor site such as dofetilide are much slower and require ~10 min to reach equilibrium (Ficker, et al., 1998). Thus, the effects of Rg3 on hERG1 (and hELK1) channel gating are extremely rapid in onset, indicating that the compound must interact with an extracellular accessible site on these channels.

**Mutation of Ser631 does not prevent effects of Rg3.** Consistent with the idea that Rg3 binds to an extracellular site of the channel, it was previously reported that a point mutation of a Ser residue located just outside the selectivity filter of the outer pore region abolished the effects of Rg3 on the hERG1 channel. Specifically, the fast deactivation of S631C hERG1 channel currents was not slowed by 3 µM Rg3 and the voltage dependence of activation was unaffected by 30 µM Rg3 (Choi, et al., 2011a). This finding is puzzling because binding of Rg3 to Ser631
should result in block of the outer pore, not activation of the channel. In contrast to this previous report we found that Rg3 affected S631C hERG1 channels in a manner similar to WT hERG1.

S631C hERG1 channels were poorly expressed in oocytes and currents were of small magnitude even when oocytes were injected with up to 40 ng of cRNA or treated with 5 mM DTT (in an effort to minimize disulfide formation). In three oocytes that expressed S631C hERG1 currents, the rate of $I_{\text{tail}}$ deactivation was dramatically slowed by 3 µM Rg3 (Fig. 2A, left panel) and the $V_{0.5}$ for activation was shifted by $-14.3 \pm 1.9$ mV (Fig. 2A, right panel; $n = 3$). To further explore the potential importance of Ser631 in the action of Rg3, we also characterized S631A and G628C/S631C hERG1 channels that express well in oocytes. The rate of S631A hERG1 channel deactivation was also greatly slowed (Fig. 2B, left panel), activation was shifted by $-13.0 \pm 1.4$ mV (Fig. 2B, right panel; $n = 4$) and the maximum $I_{\text{tail-peak}}$ was increase 1.2-fold (Supplemental Table 2) by 3 µM Rg3. G628C/S631C hERG1 channels also responded to Rg3 (Fig. 2C) with slowed $I_{\text{tail}}$ deactivation and a $V_{0.5}$ for activation shift of $-9.1 \pm 1.3$ mV ($n = 5$). Note that the tail currents of this mutant channel are inward at a $V_{\text{ret}}$ of $-70$ mV because of its reduced K$^+$ ion selectivity (Smith, et al., 1996). Together these findings indicate that Ser631 is unlikely to interact with Rg3.

**Rg3 does not interact with residues previously identified as important for other hERG1 activators.** Site-directed mutagenesis was previously used to identify specific residues that interact with several hERG1 activators. For example, L553A hERG1 channels were shown to be insensitive to RPR-260243 (Perry, et al., 2007). However, this mutation did not alter the response of hERG1 to Rg3. $I_{\text{tail}}$ deactivation was greatly slowed (Fig. 3A) and $V_{0.5}$ for activation was shifted by $-12.5 \pm 1.6$ mV (Fig. 3B; $n = 3$). Two other mutations (F557L, N658A) also prevented the slowing of deactivation of hERG1 channels by RPR-260243 (Perry, et al., 2007).
F557L and L622C abolished the effects of ICA-105574 on inactivation and deactivation (Garg, et al., 2011) and F619A prevented the effects of PD-118057 (Perry, et al., 2009). None of these five point mutations affected the ability of 3 µM Rg3 to slow channel deactivation or shift $V_{0.5}$ (Fig. 3C). Thus, it is unlikely that Rg3 binds to the same regions of the hERG1 channel previously characterized for other activators.

**Ala scan of the extracellular regions of the VSD in hERG1.** Because the onset of Rg3 is extremely fast, we reasoned that it likely binds to a region of the channel that is readily accessible from the extracellular side of the membrane. Moreover, because Rg3 affects voltage dependence of activation, we initially focused our Ala scan on the outer region of the VSD. First, we probed S3 and S4 segments and the interconnecting S3-S4 linker to search for residues that might interact with Rg3. Fig. 4A is a plot of the average shift in $V_{0.5}$ for activation and the fold-change in $\tau_{\text{deact}}$ induced by 3 µM Rg3 for mutant channels containing a single Ala substitution of 21 consecutive residues ranging from Phe508 to Arg528. Two mutations, I521A and K525A, prevented the negative shift in $V_{0.5}$ and reduced the slowing of channel deactivation compared to WT channels. Moreover, 10 µM Rg3 induced a 12.2 ± 1.1 mV ($n = 5$) positive shift in $V_{0.5}$ for activation of I521A channels, an effect that also reduced the magnitude of outward currents during depolarizing test pulses (Fig. 4B and 4C). The gating of K525A hERG1 channels was quite abnormal. These channels activated at more negative potentials ($V_{0.5} = -78.3 ± 4.0$ mV; $n = 8$) than WT channels and did not fully close at membrane voltages as negative as −120 mV (Fig. 4D). Similar to I521A channels, the voltage dependence of K525A hERG1 channel activation was shifted in the positive direction (15.2 ± 5.6 mV; $n = 7$) by 10 µM Rg3 (Fig. 4E). In summary, mutation of two residues, Ile521 and Lys525 located in the outer region of the S4
segment greatly attenuated the response to 3 µM Rg3 and responded in an unusual fashion (positive shift in $V_{0.5}$ for activation) to 10 µM Rg3.

We next extended the Ala scan to the S1-S2 extracellular region of the VSD in hERG1. Not all residues in this extensive region were probed by single mutations. Native Ala residues were not mutated to any other residue and Pro and Gly residues were not mutated since these residues are often important for secondary structure. The specific residues that were mutated to Ala are indicated in Fig. 5A. In the outer region of the S1 segment, 11 of 15 residues were mutated to Ala. In the S1-S2 linker, 13 of the 19 residues were mutated, either singly to Ala or to two different combinations of residues as indicated in the legend for Fig. 5A. In the outer region of the S2 segment, 5 of 7 residues were mutated to Ala. In addition, three other key gating residues in S2 were mutated to Ala, including Asp460 and Asp466 that form charge pairing interactions with basic residues in the S4, and Phe463 that together with Asp466 and Asp501 (in S3) form a charge-transfer center in the VSD (Tao, et al., 2010, Whicher and MacKinnon, 2016).

A plot of the average shift in $V_{0.5}$ for activation and the fold-change in $\tau_{deact}$ induced by 3 µM Rg3 for S1-S2 mutant channels are summarized in Fig. 5B. Notably, combined neutralization of four Glu residues plus two Cys to Ala mutations (E435A/E437A/E438/E444A/C445A/C449A) did not appreciably change gating properties of the channel (Fernandez, et al., 2005) or its response to Rg3. To further probe the potential importance of the acidic charges in the S1-S2 linker, three of the Glu residues and one Pro residue were all mutated to Arg (E435R/E437R/E438R/P440R). This mutant channel also responded normally to Rg3. These findings strongly suggest that Rg3 does not interact with residues in the S1-S2 linker of hERG1. However, four mutations, Y420A and V423A in the S1 segment, and L452A and F463A in the S2 segment, attenuated both the negative shift in $V_{0.5}$ and reduced the slowing of channel.
deactivation compared to WT channels. Representative currents conducted by Y420A, L452A and F463A hERG1 channels under control conditions and after treatment of oocytes with 3 and 30 µM Rg3 are illustrated in Fig. 5, panels C – E and the corresponding $g/g_{\text{max}}$-V$_t$ relationships are plotted in Fig. 5, panels F – H. At 3 µM and 30 µM, Rg3 caused a $-2.0$ mV and $-9.6$ mV shift in the $V_{0.5}$ for activation, but did not alter current magnitude of Y420A hERG1 channels (Fig. 5F). Rg3 enhanced $I_{\text{tail-peak}}$ by 25-30% and shifted the $V_{0.5}$ for activation of L452A hERG1 channels by $-6.1$ mV and $-7.3$ mV at 3 and 30 µM, respectively (Fig. 5G). F463A hERG1 channels deactivated much faster than normal and Rg3 increased $I_{\text{tail-peak}}$ by 84% and shifted the $V_{0.5}$ for activation by $-3.5$ mV at 3 µM. At 30 µM, Rg3 increased $I_{\text{tail-peak}}$ by 41%, decreased peak outward current during test pulses by $\sim50\%$ and shifted $V_{0.5}$ in a positive direction by $4.7$ mV (Fig. 5H).

Mutation of Leu417 within the S1 segment resulted in a particularly enhanced response to Rg3 (Fig. 6A). Although L417A channels deactivated very rapidly under control conditions, Rg3 still induced a dramatic slowing of this gating process (Fig. 6B). At a $V_{\text{ret}}$ of $-70$ mV, $\tau_{\text{deact}}$ was increased by a factor of $16.1 \pm 3.8$ ($n=5$), more than 3-times that observed for WT channels. Rg3 also shifted the $V_{0.5}$ for activation by $-37.4 \pm 9.2$ mV ($n=5$; Fig. 6C), 4-times that observed for WT channels.

**Ala scan of the extracellular region of the pore domain.** We next probed the outer portion of the S5 segment and the adjoining, extracellular turret to search for residues that might interact with Rg3. Supplemental Figure 1 is a plot of the average shift in $V_{0.5}$ for activation and the fold-change in $\tau_{\text{deact}}$ induced by 3 µM Rg3 for mutant channels containing a single substitution in 25 of the 45 residues between Ile567 and Tyr611. We did not mutate the single Pro or 7 Gly residues in this region and several mutant channels (K595C, Y597C, S599C) did not express.
Although not a complete scan, none of the mutant channels examined had a reduced response to Rg3 compared to WT channels.

**Effect of mutations on rate of current deactivation.** It is important to note that there was no correlation between deactivation rate of multiple mutant channels under control conditions and the fold change in $\tau_{\text{deact}}$ caused by 3 µM Rg3 (Supplemental Fig. 2). Mutations that induced fast deactivation had a variable response to Rg3. For example, D509A and L510A channels deactivated very rapidly ($\tau_{\text{deact}} = 37$ and 31 ms at −70 mV, respectively) and the Rg3 induced slowing of these channels was about half that observed for WT channels. However, other mutations that induced fast deactivation did not alter (e.g., D460A, $\tau_{\text{deact}} = 67$ ms; D466A, $\tau_{\text{deact}} = 59$ ms), or even increased (e.g., L417A, $\tau_{\text{deact}} = 43$ ms; Y569A, $\tau_{\text{deact}} = 104$ ms; N658A, $\tau_{\text{deact}} = 128$ ms) the fold change in $\tau_{\text{deact}}$ induced by Rg3.

**Double mutations nearly eliminate the action of Rg3.** We next paired Y420A with either L452A or I521A into a single construct. Each of these single mutations reduced the response to Rg3 (Figs. 4 and 5). Y420A/L452A hERG1 channels activated at more positive potentials ($V_{0.5} = 5.2 \pm 6.7$ mV, $n = 14$) and deactivated more rapidly than WT channels (Fig. 7A). The double mutant channel current also displayed an unusual tendency to decrease in magnitude (“run-down”) during repetitive pulsing soon after impaling an oocyte with recording microelectrodes. Run-down typically reduced currents to about half their initial amplitude before reaching a steady-state level. Currents for this mutant channel were not recorded until after run-down was complete. Rg3 at 3 and 30 µM increased $I_{\text{peak}}$ and $I_{\text{tail-peak}}$ for $V_t > 0$ mV, but did not significantly alter the voltage dependence of Y420A/L452A hERG1 channel activation (Fig. 7A-C). $V_{0.5}$ for activation was shifted by $2.4 \pm 4.9$ mV ($n = 14$) at 3 µM Rg3 and $-1.3 \pm 3.6$ mV ($n = 9$) at 30 µM Rg3 (Fig. 7B).
Y420A/L452A hERG1 channels were fully activated by applying a prepulse to 40 mV for 1 s before repolarization to a $V_{ret}$ that was varied from 20 mV to −130 mV in 10-mV increments (Fig. 7D). For a $V_{ret}$ of −40 to −130 mV, deactivation of current was sufficient for accurate kinetic measurements. Rg3 had only minimal effects on the fully-activated $I_{tail}$-$V_t$ relationship (Fig. 7E). Rg3 at 3 µM slightly slowed (by 1.14-fold at −70 mV) the rate of $I_{tail}$ deactivation, whereas 30 µM Rg3 had no effect on $\tau_{deact}$ (Fig. 7F). Rg3 had little or no effect on the relative amplitude of the slowly deactivating component of $I_{tail}$, including at return potentials (i.e., −40 mV) where $\tau_{deact}$ was slowest (Fig. 7G). By contrast, 10 µM Rg3 increased $\tau_f$ and $\tau_s$ for WT hERG1 channel deactivation at a $V_{ret}$ of −70 mV by 6-fold and 11-fold, respectively and increased the relative amplitude of the slowly deactivating component of $I_{tail}$ (Wu, et al., 2016). The negligible effects of Rg3 on $\tau_{deact}$ of Y420A/L452A hERG1 channels were not simply due to the fact that these mutant channels deactivate rapidly in the absence of Rg3. As shown in Supplemental Figure 2, there is no correlation between the control $\tau_{deact}$ and the fold change in this parameter induced by Rg3.

Y420A/I521A hERG1 channel currents (Fig. 8A) activated at more negative potentials ($V_{0.5} = −38.2 ± 4.2$, $n = 6$) than WT channels. In response to 3 and 30 µM Rg3, $V_{0.5}$ was shifted by −1.8 ± 1.5 mV ($n = 6$) and 8.2 ± 0.9 mV ($n = 3$), respectively (Fig. 8B). The maximum $I_{tail}$-$V_t$ for Y420A/I521A hERG1 channels was increased by 3 µM Rg3 (Fig. 8B), while 30 µM Rg3 caused a reduction in $I_{peak}$ (Fig. 8C). Although Rg3 had almost no effect on $V_{0.5}$ or $\tau_{deact}$ of Y420A/L452A channels, it increased the maximum value of $I_{tail}$-$V_t$ by a factor of −1.3 (Supplemental Table 2). For comparison, the fold change in $I_{tail}$-$V_t$ was 0.9 to 1.2 for the majority of other mutant channels and 1.1 for WT channels (Supplemental Figure 3). The relatively large increase in $I_{tail}$-$V_t$ observed for Y420A/L452A channels may be related to the
fast rate of deactivation for these channels under control conditions, especially considering that
the fold increase in $I_{\text{tail-peak}}$ was 1.02 and 1.23 for the single mutant (Y420A or L452A) hERG1
channels, respectively. The increase in $I_{\text{tail-peak}}$ was > 1.6-fold in most other fast deactivating
mutant channels, including D456A, F463A, D509A, L510A and F619A hERG1 (Supplemental
Figure 3). We suspect that $I_{\text{tail-peak}}$ for these channels may have been underestimated under
control conditions because deactivation was too fast to permit adequate recovery of channels
from an inactivated state. Another possibility is that Rg3 activates channels by multiple
mechanisms, possibly by binding to multiple sites including one that is not altered by the double
mutation Y420A/L452A. Finally, the reduced efficacy of Rg3 on the gating of the double mutant
channels is not simply the sum of the effects of single mutations (Fig. 8D and 8E). The
combination of Y420A and L452A mutations were most effective in suppressing the action of
Rg3.

In summary, mutation to Ala of Leu417 and Tyr420 in the S1 segment, Leu452 and Phe463 in
the S2 segment, and Ile521 and Lys525 in the S4 segment produced the most dramatic changes
in the effects of Rg3 on hERG1 channel gating. These findings suggest that these residues are
either structural determinants of the Rg3 binding site or that mutation of these particular residues
alters the response to Rg3 by an indirect, allosteric mechanism. We employed molecular
modeling to help distinguish between these possibilities.

Docking and MD simulations suggest that the VSD of hERG1 can accommodate Rg3.
Simulated docking of Rg3 to molecular models of the hERG1 voltage sensor module in two
different activated states was performed. It is not clear if the VSD in the rEAG1 structure
template represents a fully activated state (Whicher and MacKinnon, 2016), since only the first
three positive charges are above the charge transfer center. Thus, we additionally built a model,
where the alignment of helix S4 was shifted upward by three residues (denoted activated₁), to obtain a model with four positive charges above the charge transfer center. This activated₁ model is consistent with experimental work from Zhang, et al. (2005), suggesting a direct interaction between residues D456 from helix S2 and Arg528 from the S4 segment. In both models, docking predicts that Rg3 can protrude deeply into the VSD modules (Fig. 9A-D and Supplemental Figure 4), where they remain stably bound during 50 ns of unbiased MD simulations (Supplemental Figure 5 A,B). Detailed views of the putative drug-binding regions in the activated and activated₁ states are depicted in Fig. 9, in which residues identified by scanning mutagenesis and a few other residues in close contact (within 4 Å) with the drug molecule are shown in stick representation. In both activated states residues Tyr420, Thr421, Leu452, Asp460, Asp509, Ile512, Ile521, Gly522 and Lys525 and Arg528 are within 4 Å of the Rg3 molecule. A second class of high impact residues (as determined by mutagenesis) is not in direct contact with Rg3 in either model; however, a downward movement of the drug is already apparent in the activated model, where the distance between Rg3 and Phe463 is approaching 3.5 Å, compared to >5 Å in the activated₁ model (measured at the end of the 50 ns runs). Thus it is plausible that residues Phe463, Val418 and Leu417 might come in close contact with the drug upon voltage sensor downward movement. The 3rd group of residues is unlikely to form direct interactions according to our homology models, even when taking into consideration conformational rearrangements upon gating. Most notable are residues Ile419 (S1) and Leu510 (S3) that are lipid interacting, facing the opposite site of the predicted binding site. Ser428 is predicted to form a stable hydrogen bond to Rg3; however S428A hERG1 channels responded normally to Rg3. Further, loop residues including Glu435 can be within 4 Å of the drug, but due to the high flexibility of these regions (see Supplemental Figure 6) might not contribute to binding.
Hydrogen bond contacts between Rg3 and hERG1 in the activated and activated1 model are summarized in Supplemental Table 3. Finally, during the revision of this manuscript the Mackinnon group published the first cryo-EM structure of the hERG1 channel in the open state (Wang and MacKinnon, 2017). A structural alignment of the VSDs of hERG1 with rEAG1 reveals similar conformations in this domain with an RMSD of 1.25 Å (see Supplemental Figure 7). However, the hERG1 VSD has a lower resolution (~4 - 5Å) than the EAG1 structure and is also missing the S1-S2 and S3-S4 extracellular loops. For these reasons the new structure provided no advantage for our simulations.

Discussion

**Rg3 interacts with the VSD of hERG1.** Ginsenoside Rg3 is a large steroid glycoside that slows the rate of deactivation and shifts the voltage dependence of hERG1 channel activation to more negative potentials (Choi, et al., 2011a, Wu, et al., 2016). These effects are similar to a few other hERG1 activators, including mallotoxin (Zeng, et al., 2006) and NS1643 (Xu, et al., 2008) and related analogs (Guo, et al., 2014). However, unlike these other compounds the onset of Rg3 activity upon extracellular application to cells is extremely rapid, suggesting that it likely binds to an extracellular accessible site on the hERG1 channel. S631C hERG1 channels were reported to be insensitive to Rg3 (Choi, et al., 2011a), suggesting that the alkaloid might interact directly or indirectly with Ser631, a residue located within the outer vestibule of the channel in close proximity to its selectivity filter. However, our finding that Rg3 altered the gating of channels harboring Ser631 mutations (i.e., S631C, S631A, or G628C/S631C) in a manner similar to WT channels indicated to us that this residue does not interact with Rg3. Therefore, we mutated 84 other residues located in the extracellular facing region of the VSD to search for regions of the hERG1 subunit that may interact directly or indirectly (allosterically) with Rg3. We focused our
attention on mutations that reduced both of the primary effects of Rg3. Our scanning
mutagenesis identified 5 mutations (Y420A, L452A, F463A, I521A, K525A) that highly
diminished the slowing of deactivation and the negative shift in $V_{0.5}$ for activation normally
induced by Rg3 and one mutation (L417A) that greatly enhanced these effects of Rg3. In
addition, mutation of 11 additional residues enhanced either the ability of Rg3 to slow
deactivation or shift $V_{0.5}$ for activation and 4 other mutations caused a reduced deactivation
response. The combined point mutations of Y420A and L452A abolished the effects of Rg3 on
$V_{0.5}$ or $\tau_{\text{deact}}$ at the high concentration of 30 µM. The double mutation also induced fast channel
deactivation ($\tau_{\text{deact}} = 73$ ms at −70 mV), but this does not explain why Rg3 did not slow
deactivation as illustrated in Supplemental Figure 2 that shows no correlation between
deactivation rate of multiple mutant channels under control conditions and the fold change in
$\tau_{\text{deact}}$ caused by 3 µM Rg3. The side chains of most, but not all of these “high impact” residues
face towards a cleft located between the transmembrane segments of the VSD. Simulated
molecular docking of Rg3 to a homology model of the hERG1 VSD based on the cryo-EM
structure of the rat EAG1 channel (Whicher and MacKinnon, 2016) corroborated some, but not
all of the scanning mutagenesis findings. For example, a stable H-bond was predicted between
Ser428 and Rg3, but the S428A mutant channel responded normally to Rg3. The residue
interactions predicted by the docking model were primarily hydrophobic in nature and the
binding mode for Rg3 was very stable with an RMSD value of ~0.15 nm during repeated MD
simulations of 50 nanoseconds. Together these findings suggest that Rg3 modifies the gating of
hERG1 channels by binding to the outer region of the VSD. Presumably this binding mode
favors the activated state of the VSD and shifts the $V_{0.5}$ for activation to more negative potentials.
Further studies are required to determine if slow deactivation results from a slower rate of
repolarization induced S4 charge displacement or from a functional uncoupling between S4 movement and closure of the activation gate formed by the S6 bundle crossing. We previously found that the profound slowing of hERG1 deactivation caused by RPR-260243 (RPR) was not associated with a change in the kinetics of gating current. The putative binding site for RPR is located on the pore domain, not in the VSD as proposed here for Rg3. Further evidence that RPR and Rg3 do not interact at a common site on hERG1 was our finding that mutations that drastically alter the response to RPR (i.e., L553A, F557L, N658A) did not alter the response to Rg3. Thus, it is uncertain whether the molecular mechanism of slowed channel deactivation caused by RPR and Rg3 is similar or entirely different.

Limitations of study. We used a site-directed mutagenesis approach to identify the molecular determinants of Rg3-induced changes in hERG1 channel gating. This indirect approach of identifying the putative binding site for a ligand has several important limitations. Perhaps the most significant is the underlying assumption that an altered response to Rg3 induced by a single or multiple point mutations can be used to identify specific residues that interact with Rg3. It is not possible to determine if an altered response to Rg3 induced by a point mutation results directly from a decreased binding affinity or from an indirect, allosteric mechanism unrelated to a change in binding affinity. For example, while mutation to Ala of Ile419 in the S1 segment and Leu510/Leu511 in the S3 segment altered the response of hERG1 to Rg3, these residues are not predicted to make contact with the alkaloid in the dockings shown in Fig. 9. A second limitation is that many mutations significantly alter the properties of hERG1 channel gating (Supplemental Tables 1 and 2), perhaps in a manner that would limit or accentuate the effects of Rg3. However, of the 18 mutations that shifted the $V_{0.5}$ for activation by $> +10 \text{ mV}$, only 3 mutations (Y420A, L452A, F463A) showed a reduced $V_{0.5}$ shift in response to Rg3. Of the 8 mutations that shifted
V_{0.5} to a more negative potential by > −10 mV compared to WT channels, only 3 mutations (I521A, G522A, K525A) exhibited a reduced V_{0.5} shift in response to Rg3. Many mutations also altered the rate of channel deactivation. However, there was no correlation between $\tau_{\text{deact}}$ measured under control conditions and the fold-change in this parameter induced by Rg3 (Supplemental Figure 2). Thus, although many mutations significantly altered channel gating properties, these changes cannot account for the finding that specific mutations altered the response to Rg3. These findings argue against the proposal that activators must “work harder to overcome mutation induced loss of baseline channel function” (Pearlstein, et al., 2017). Third, for some mutant channels, fitting $I_{\text{tail}}$ deactivation with a single exponential function led to an underestimation of the fold change in $\tau_{\text{deact}}$. This is the case for WT channels where Rg3 increases the $\tau$ for the fast and slow components of deactivation by 5-fold and 10-fold, respectively (Wu, et al., 2016). Finally, most residues were only mutated to Ala and not all residues were mutated. Mutation to a residue other than Ala may have identified additional native residues as structural determinants of Rg3 activity. We also chose not to mutate most native Ala, Pro or Gly residues (e.g., in S1-S2 region) and therefore, would have missed if any of these residues interact with Rg3.

**Therapeutic implications.** Activators of hERG1 channels have been proposed as a pharmacological approach for prevention of arrhythmia associated with congenital or acquired LQTS (Kang, et al., 2005, Sanguinetti, 2014, Wu and Sanguinetti, 2016). However, excessive activation of hERG1 would cause early repolarization of action potentials, essentially mimicking congenital short QT syndrome, a disorder of ventricular repolarization that also increases the risk of life-threatening arrhythmia (Gaita, et al., 2003, Shah, 2010). Rg3 and related ginsenosides have been extensively studied and are claimed to exert a multitude of therapeutic effects (Ahn, et
al., 2016, Nah, 2014, Weber, et al., 2012). Given the plethora of activities associated with Rg3, it is not surprising that this ginsenoside is not a hERG1 specific compound. Rg3 also inhibits Na⁺, 5-HT3A and nicotinic acetylcholine receptor channels (Jeong, et al., 2004), L-type Ca²⁺ (Choi, et al., 2009) and KCNQ1 channels (Choi, et al., 2010). Rg3 modifies gating to activate KCNQ1/KCNE1 (Choi, et al., 2010), BKCa (Choi, et al., 2011b) as well as EAG1, ERG3 and ELK1 channels (Wu, et al., 2016). Some of these channel activities occur over a range of concentrations similar to those that affect hERG1 channels. Thus, while Rg3 provides a structural starting point for design of novel compounds that can activate hERG1, its nonspecific activities makes it an inappropriate agent for evaluating the efficacy or safety of hERG1 activators in LQTS.

Other compounds that interact with the VSD of voltage gated K⁺ or Na⁺ channels have been reported to activate or inhibit channels in a specific manner. For example, ICA-27243 (N-(6-chloro-pyridin-3-yl)-3,4-difluoro-benzamide) selectively binds to the VSD and shifts the voltage dependence of KCNQ2/3 (Kv7.2/.3) channels to more negative potentials (Padilla, et al., 2009). GX-936 and related aryl sulfonamides selectively inhibit Nav1.7 channels by binding to, and trapping the 4th VSD in its activated state. This action prevents deactivation of the VSD and thereby stabilizes the inactivated state of the Nav1.7 channel (Ahuja, et al., 2015). Future studies are required to determine if structural modifications of Rg3 might result in a compound that specifically interacts with the VSD of hERG1 channels.

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Authorship Contributions

Participated in research design: Gardner, Sanguinetti.

Conducted experiments: Gardner, Wu, Thomson, Sanguinetti.

Performed data analysis: Gardner, Sanguinetti.

Performed molecular modeling: Zangerl-Plessl, Stary-Weinzinger.

Wrote or contributed to the writing of the manuscript: Sanguinetti, Stary-Weinzinger.
References


Footnotes

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Figure Legends

**Fig. 1.** Ginsenoside Rg3 alters activation gating of hERG1 channels. (A) Voltage clamp pulse protocol (upper left panel) and whole cell currents (bottom panels) recorded from a single oocyte expressing WT hERG1 before (control) and after treatment with 3 µM Rg3. Small arrows indicate zero current level. (B) Rg3 shifts the $I_{\text{peak}}$-$V_t$ relationship for hERG1 to more negative potentials. (C) Rg3 shifts the voltage dependence of hERG1 activation to more negative potentials. Peak $I_{\text{tail}}$ at $-70$ mV from currents shown in panel A were plotted as a function of $V_t$ and fitted with a Boltzmann function (smooth curves). For control, $V_{0.5} = -31.5$ mV, $k = 8.1$; for 3 µM Rg3, $V_{0.5} = -42.6$ mV, $k = 9.9$. (D) Currents recorded in response to repetitive pulses, applied once every 5 s to the indicated voltage step shown in upper panel, before (0) and during the 1st, 2nd and 3rd steps (1, 2, 3) after start of bath perfusion of oocyte chamber with a solution containing 3 µM Rg3. (E) Onset of Rg3 action is rapid. $I_{\text{tail}}$ magnitude (measured at 1 s after return of voltage to $-70$ mV) was normalized to the peak value for each oocyte, plotted as a function of time after start of Rg3 application, then fitted with a single exponential function (dashed curve). The time constant for onset of Rg3 action was $1.9 \pm 1.1$ s ($n = 7$).

**Fig. 2.** Ser631 is not a structural determinant of the Rg3 binding site. (A-C) Left panels show representative current traces recorded during 4 s pulses to 0 mV followed by 4 s pulse to $-70$ mV for oocytes expressing S631C hERG1 (A), S631A hERG1 (B) or G628C/S631C hERG1 (C) mutant channels. Note that 3 µM Rg3 slows the rate of current deactivation at $-70$ mV for all channels. Right panels are plots of $g/g_{\text{max}}$-$V_t$ relationships for indicated mutant hERG1 channel types. Data were normalized to $g_{\text{max}}$ determined under control conditions and fitted with a Boltzmann function (smooth curves). $V_{0.5}$ and $k$ values for all fits are presented in Supplemental
Table 1. The shift in $V_{0.5}$ induced by Rg3 was $-14.3 \pm 1.9$ mV for S631C ($n = 3$), $-12.7 \pm 1.4$ mV for S631A ($n = 4$) and $-9.1 \pm 1.3$ mV for G628C/S631C ($n = 5$) hERG1 channels.

**Fig. 3.** Rg3 does not interact with hERG1 residues important for activity of other activators. (A) Voltage clamp pulse protocol (upper left panel) and whole cell currents recorded from an oocyte expressing L553A hERG1 channels before (control) and after treatment with 3 µM Rg3. Small arrows indicate zero current level. (B) $g/g_{\text{max}} - V_t$ relationship for L553A hERG1 channels. Data were normalized to $g_{\text{max}}$ determined under control conditions and fitted with a Boltzmann function (smooth curves). $V_{0.5}$ and $k$ values for fits are presented in Supplemental Table 1. The shift in $V_{0.5}$ induced by Rg3 was $-12.5 \pm 1.6$ mV ($n = 3$). (C) Plot of shift in $V_{0.5}$ for activation (black bars) and fold-change in $\tau_{\text{deact}}$ (grey bars) induced by 3 µM Rg3 for WT or mutant hERG1 channels as indicated on x-axis ($n$ indicated below $V_{0.5}$ bars).

**Fig. 4.** Ala scan of select residues located in the S3-S4 region of hERG1. (A) Plot of shift in $V_{0.5}$ for activation (black bars) and fold-change in $\tau_{\text{deact}}$ (grey bars) induced by 3 µM Rg3 for WT or mutant hERG1 channels as indicated on x-axis ($n$ indicated below $V_{0.5}$ bars). Point mutations were introduced into the S3 segment, S3-S4 linker (S34L) or S4 segment. (B) Voltage clamp pulse protocol (upper panel) and whole cell currents recorded from an oocyte expressing I521A hERG1 channels before (control) and after treatment with 10 µM Rg3. Small arrows indicate zero current level. (C) Plot of $g/g_{\text{max}} - V_t$ relationships for I521A hERG1 channels. Data were normalized to $g_{\text{max}}$ determined under control conditions and fitted with a Boltzmann function (smooth curves). $V_{0.5}$ and $k$ values for fits are presented in Supplemental Table 1. Rg3 shifted $V_{0.5}$ of I521A hERG1 channels by $2 \pm 2.0$ mV at 3 µM ($n = 8$) and by $12.2 \pm 1.1$ mV at 10 µM ($n = 5$). (D) Whole cell currents recorded from an oocyte expressing K525A hERG1 channels before (control) and after treatment with 10 µM Rg3. Small arrows indicate zero current level. (E) Plot
of \( g/g_{\text{max}} - V_t \) relationships for K525A hERG1 channels. Data were normalized to \( g_{\text{max}} \) determined under control conditions and fitted with a Boltzmann function (smooth curves). Rg3 shifted \( V_{0.5} \) of K525A hERG1 channels by \( 1.1 \pm 2.7 \text{ mV} \) at \( 3 \mu M \) \((n = 5)\) and by \( -0.2 \pm 5.1 \text{ mV} \) at \( 10 \mu M \) \((n = 8)\). \( V_{0.5} \) and \( k \) values for Boltzmann fits in panels C and E are presented in Supplemental Table 1.

**Fig. 5.** Ala scan of select residues located in the S1-S2 region of hERG1. (A) Amino acid sequence (single letter code) of extracellular regions of S1 and S2 segments and S1-S2 extracellular linker (S12L). Single residues that were mutated to Ala are shown in bold text. Underlined residues designate the multiple residues that were mutated to construct 4E2C/6A (E435A/E437A/E438A/E444A/C445A/C449A) or 3E1P/4R (E435R/E437R/E438R/P440R) multi-mutation channels. (B) Plot of shift in \( V_{0.5} \) for activation (black bars) and fold-change in \( \tau_{\text{deact}} \) (grey bars) induced by 3 \( \mu M \) Rg3 for WT or mutant hERG1 channels as indicated on x-axis \((n \text{ indicated below } V_{0.5} \text{ bars})\). Point mutations were introduced into the S1 segment, S1-S2 linker \((S12L)\) or S2 segment. 4E2C/6A = E435A/E437A/E438/E444A/C445A/C449A (Fernandez, et al., 2005); 3E1P/4R = E435R/E437R/E438R/P440R. (C-E) Voltage clamp pulse protocol (upper left panel) and whole cell currents recorded from an oocyte expressing Y420A (C), L452A (D) and F463A (E) hERG1 channels before (control) and after treatment with 3 and 30 \( \mu M \) Rg3. Small arrows indicate zero current level. (F-H) Plot of \( g/g_{\text{max}} - V_t \) relationships for Y420A (F), L452A (G) and F463A (H) hERG1 channels. Data were normalized to \( g_{\text{max}} \) determined under control conditions and fitted with a Boltzmann function (smooth curves). \( V_{0.5} \) and \( k \) values for fits are presented in Supplemental Table 1. Rg3 shifted \( V_{0.5} \) of Y420A hERG1 channels by \( -2 \pm 2.4 \text{ mV} \) at \( 3 \mu M \) \((n = 9)\) and by \( -9.6 \pm 1.9 \text{ mV} \) at \( 30 \mu M \) \((n = 7)\). Rg3 shifted \( V_{0.5} \) of L452A hERG1 channels by \( -6.1 \pm 1.2 \text{ mV} \) at \( 3 \mu M \) \((n = 4)\) and by \( -7.3 \pm 0.9 \text{ mV} \) at \( 30 \mu M \) \((n = 3)\). Rg3
shifted $V_{0.5}$ of F463A hERG1 channels by $-3.5 \pm 2.4$ mV at 3 µM ($n = 7$) and by $4.7 \pm 2.6$ mV at 30 µM ($n = 4$).

**Fig. 6.** Rg3 dramatically slows deactivation and causes a large negative shift in the voltage dependence of activation of L417A hERG1 channels. (A) Voltage clamp pulse protocol (upper left panel) and whole cell currents recorded from an oocyte expressing WT hERG1 before (control) and after treatment with 3 µM Rg3. Small arrows indicate zero current level. (B) Rg3 slows rate of $I_{\text{tail}}$ deactivation measured at a $V_{\text{ret}}$ of $-70$ mV. In 7 oocytes, $\tau_{\text{deact}}$ before and after treatment with 3 µM Rg3 was 43 ± 13 ms and 696 ± 336 ms, respectively. (C) $g/g_{\text{max}}$-$V_t$ relationship for L417A hERG1 channels. Data were normalized to $g_{\text{max}}$ determined under control conditions and fitted with a Boltzmann function (smooth curves). For control conditions: $V_{0.5} = -5.7 \pm 6.9$ mV, $k = 12.2 \pm 1.1$; after 3 µM Rg3: $V_{0.5} = -43.1 \pm 7.8$ mV, $k = 5.03 \pm 0.42$ ($n = 5$). The shift in $V_{0.5}$ induced by Rg3 was $-37.4 \pm 9.2$ mV ($n = 5$).

**Fig. 7.** Effect of Rg3 on Y420A/L452A hERG1 channel currents. (A) Voltage clamp pulse protocol (upper left panel) and whole cell currents recorded from an oocyte expressing Y420A/L452A hERG1 channels before (control) and after treatment with 3 and 30 µM Rg3. Small arrows indicate zero current level. (B) Plot of $g/g_{\text{max}}$-$V_t$ relationship for Y420A/L452A hERG1 channels. Data were normalized to $g_{\text{max}}$ determined under control conditions and fitted with a Boltzmann function (smooth curves). $V_{0.5}$ and $k$ values are presented in Supplemental Table 1. Rg3 shifted $V_{0.5}$ by $2.4 \pm 4.9$ mV at 3 µM ($n = 14$) and $-1.3 \pm 3.6$ mV at 30 µM ($n = 9$). (C) $I_{\text{peak}}$-$V_t$ relationship for Y420A/L452A hERG1 channels. Data were normalized to the maximum $I_{\text{peak}}$ determined under control conditions. (D-G) Rg3 has only minor effects on voltage-dependent gating of Y420A/L452A hERG1 channels. (D) Voltage clamp pulse protocol (upper left panel) and tail currents recorded from an oocyte expressing Y420A/L452A hERG1
channels before (control) and after treatment with 3 and 30 µM Rg3. For these experiments the holding potential was −70 mV and tail currents were measured at the indicated $V_{\text{ret}}$ following a 1 s prepulse to 40 mV. Pulses were applied once every 12 s. Currents recorded at a $V_{\text{ret}}$ ranging from 20 mV to −30 mV are indicated by dashed curves. Currents recorded at a $V_{\text{ret}}$ ranging from −40 to −130 mV are indicated by solid curves. (E) Plot of average normalized $I_{\text{tail-peak}}$ vs $V_{\text{ret}}$ before (control) and after treatment of oocytes with 3 and 30 µM Rg3. (F) Effect of Rg3 on the two time constants of deactivation of hERG1 current deactivation as a function of $V_{\text{ret}}$. (G) Effect of Rg3 on the relative magnitude of the slow component of current deactivation as a function of $V_{\text{ret}}$. For panels E-G, $n = 8$.

**Fig. 8.** Effect of Rg3 on Y420A/I521A hERG1 channel currents. (A) Voltage clamp pulse protocol (upper left panel) and whole cell currents recorded from an oocyte expressing Y420A/I521A hERG1 channels before (control) and after treatment with 3 and 30 µM Rg3. (B) $g/g_{\text{max}}-V_t$ relationships for Y420A/I521A hERG1 channels. $V_{0.5}$ and $k$ values are presented in Supplemental Table 1. Rg3 shifted $V_{0.5}$ by $−1.8 \pm 1.5$ mV at 3 µM ($n = 6$) and by $8.2 \pm 0.9$ mV at 30 µM ($n = 3$). (C) $I_{\text{peak}}-V_t$ relationship for Y420A/I521A hERG1 channels. Data were normalized to the maximum $I_{\text{peak}}$ determined under control conditions ($n$ was same as indicated in panel B). (D) Plot of shift in $V_{0.5}$ for activation induced by 3 µM Rg3 (black bars), 10 µM Rg3 (*control, grey bars) or 30 µM Rg3 (mutants, grey bars) for indicated WT or mutant channels ($n$ indicated for each bar). (E) Plot of fold-change in $\tau_{\text{deact}}$ induced by 3 µM Rg3 (black bars), 10 µM Rg3 (*control, grey bars) or 30 µM Rg3 (grey bars) for indicated WT or mutant hERG1 channels ($n$ was same as indicated in panel D).
Fig. 9. Docking interactions of Rg3 with the voltage sensor models in two activated state models. (A, B) rEAG1 based activated VSD module of the hERG1 channel shown in side view. Rg3 is shown in green stick representation. Key residues identified in scanning mutagenesis are labeled in bold text and shown in stick representation. Additional residues shown in plain text (e.g., S428, F424, E435, D456) are also predicted to be in close contact (within 4 Å) with Rg3, but exhibited normal drug response when mutated to Ala. (C, D) Activated VSD module with residue labeling the same as in panels A and B.
Figure 8
Figure 9