Ranolazine inhibition of hERG potassium channels: Drug–pore interactions and reduced potency against inactivation mutants

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Abstract

The antianginal drug ranolazine, which combines inhibitory actions on rapid and sustained sodium currents with inhibition of the hERG/Ko potassium channel, shows promise as an antiarrhythmic agent. This study investigated the structural basis of hERG block by ranolazine, with lidocaine used as a low potency, structurally similar comparator. Recordings of hERG current (IhERG) were made from cell lines expressing wild-type (WT) or mutant hERG channels. Docking simulations were performed using homology models built on MthK and KvAP templates. In conventional voltage clamp, ranolazine inhibited IhERG with an IC50 of 8.03 μM; peak IhERG during ventricular action potential clamp was inhibited ~62% at 10 μM. The IC50 values for ranolazine inhibition of the S620T inactivation deficient and N588K attenuated inactivation mutants were respectively ~73-fold and ~15-fold that for WT IhERG. Mutations near the bottom of the selectivity filter (V625A, S624A, T623A) exhibited IC50s between ~8 and 19-fold that for WT IhERG, whilst the Y652A and F656A S6 mutations had IC50s ~22-fold and 53-fold WT controls. Low potency lidocaine was comparatively insensitive to both pore helix and S6 mutations, but was sensitive to direction of K+ flux and particularly to loss of inactivation, with an IC50 for S620T-hERG ~49-fold that for WT ihERG. Docking simulations indicated that the larger size of ranolazine gives it potential for a greater range of interactions with hERG pore side chains compared to lidocaine, in particular enabling interaction of its two aromatic groups with hERG pore side chains of both Y652 and F656. The N588K mutation is responsible for the SQT1 variant of short QT syndrome and associated Torsades de Pointes (TdP) arrhythmia [12,13].

1. Introduction

Class III antiarrhythmic agents prolong refractoriness and can be effective in treating atrial and ventricular re-entrant arrhythmias [1,2]. A number of ‘pure’ Class III compounds, including β-sotalol and dofetilide exert their effects through inhibition of the rapid delayed rectifier K+ current, IKr, and its recombinant equivalent hERG [3,4]. IKr/hERG blockade additionally contributes to the actions of the Class Ia antiarrhythmics quinidine and disopyramide and has also been reported for Class Ic drugs such as propafenone and flecaïnide (e.g.[5–8]). By contrast, the Class Ib antiarrhythmic agents mexiletine and lidocaine have comparatively low affinity for Ik/hERG channels [9–11]. Deleterious as well as beneficial effects of Class Ia and III drugs are associated with their IKr/hERG blocking propensity, as they carry some risk of acquired long QT syndrome and associated Torsades de Pointes (TdP) arrhythmia [12,13].

The antianginal agent ranolazine is receiving increasing interest as a potential antiarrhythmic agent [14,15]. In a pre-clinical experimental setting ranolazine can produce modest prolongation of action potential duration (APD; without reverse-rate dependence), without augmenting transmural dispersion of repolarization and it can suppress early afterdepolarizations in Purkinje fibres and ventricular M’ cells; thus ranolazine appears to be without the potential to produce TdP of pure Class III drugs (reviewed in [16, 17]). Patients with angina receiving ranolazine exhibit concentration-dependent QT interval prolongation that is only modest at high doses (QT, increase of ~10 ms with 1000 mg twice daily; [18]). In humans, continuous ECG monitoring of 6351 patients over 7 days following admission for acute coronary syndrome (MERLIN-TIMI 36) showed that fewer patients on ranolazine had episodes of ventricular or supraventricular tachycardia, with a trend also towards a lower incidence of new onset atrial fibrillation (AF) [19]. A small-scale study of patients with new onset or paroxysmal atrial fibrillation (AF; most of whom had some structural heart disease) showed 72% (13/18 patients) conversion to sinus rhythm with high dose (2000 mg) ranolazine, suggesting that further evaluation of ranolazine for AF conversion is warranted [20]. A study of 25 patients with AF resistant to electrical cardioversion, found that 19 patients...
were successfully cardioverted within two attempts, following oral ranolazine administration [21].

Ranolazine exerts a use-dependent block of both rapid and sustained Na⁺ currents (iNa and iNa,Late respectively) [17,22,23] and due to differences between atrial and ventricular iNa, the drug is considered to exhibit atrioreselectivity of rapid iNa block [24]. At clinically relevant concentrations, ranolazine also inhibits iNaK and its recombinant equivalent ‘hERG’ [23,25,26]. Consequently, ranolazine combines clinically relevant actions against iNaK and iNa and this appears to lead to a more favourable frequency dependence of its effects on cardiac repolarization than seen with pure iNaK blockade [27]. Indeed, ranolazine has a more favourable action on atrial effective refractory period than expected for pure iNaK blockers such as dofetilide [28]. Ranolazine has been reported to produce hERG block that is rapid in its onset and reversibility and that depends on drug-access to the channel from the cell interior [26]. Additionally, whilst ranolazine does not impair trafficking of wild-type (WT) hERG channels to the cell membrane [26], a recent study suggests that it can promote trafficking of long QT syndrome (LQTS) mutant channels and thus ranolazine may have potential to act as a pharmacological chaperone for LQTS mutant hERG channels [29]. However, although there is evidence that ranolazine binds to hERG within the channel’s inner cavity [23], the nature of the underlying drug–channel interactions is incompletely understood and the role of channel inactivation (which is critical for hERG inhibition by some, but not all, drugs [8,30]) in the drug’s hERG blocking action has not been studied. Accordingly, the present investigation was undertaken to elucidate the nature and molecular determinants of the drug’s inhibition of HREK, through a combination of electrophysiology and mutagenesis, together with drug docking to homology models of the hERG channel. As noted by others [31], ranolazine shares chemical structure with lidocaine (see Fig. S1 in the online supplement); both compounds exhibit a tertiary amine local anesthetic (LA) structure containing hydrophobic (aromatic ring) and hydrophilic (tertiary amine) groups [31]. Lidocaine was therefore used as a low affinity [11], structurally similar comparator for experiments with hERG mutants and for docking.

2. Materials and methods

2.1. Maintenance of mammalian cell lines and cell transfection

HEK 293 cells stably expressing WT hERG (provided by Professor Craig January [32]) or the Y652A and F656A mutants [33] were maintained as described previously (e.g. [33,34]). For transient transfection, cells were maintained as previously described [35] and transfected with cDNA plasmids (T623A, S624A, V625A, N588K, S620T hERG) using either Lipofectamine 2000 or LTX (Invitrogen, Paisley, UK). Green Fluorescent Protein or CD8 was used as markers of successful transfection. Further details are available in the online supplementary information.

2.2. Electrophysiological recordings

Recordings were made at 37 ± 1 °C with a standard Tyrode’s solution containing (in mM): 140 NaCl, 4 KCl, 2 CaCl₂, 1 MgCl₂, 10 Glucose, 5 HEPES (titrated to pH 7.4 with NaOH). For experiments with mutants T623A and F656A and their WT controls, the superfusate contained 94 mM K⁺ (with NaCl concentration correspondingly reduced) [34]. Patch-pipettes (Glass 8250, AM Systems Inc.) had a final resistance of 2–4 Mȍ. The pipette dialysis solution contained (in mM): 130 KCl, 1 MgCl₂, 5 EGTA, 5 MgATP; pipette pH was buffered to pH 7.2 with 10 mM HEPES (titrated with KOH). Series resistance values typically lay between 3 and 7 Mȍ and were compensated by ~60–80%. The data digitization rate was 10 kHz during all protocols and a bandwidth of 2 kHz was set on the amplifier. Further information, including specific equations used for data analysis, is available in the online supplementary information.

2.3. Ranolazine and lidocaine

Ranolazine di-HCl powder (Sequoia Research Products Ltd. UK) was dissolved in deionized water (Milli-Q, Millipore) to give a stock concentration of 10 mM. Lidocaine was dissolved in deionized water at a stock concentration of 10–20 mM. Both stock solutions were diluted with appropriate superfusion solutions to achieve series of concentration in the Results section. For higher final concentrations, lidocaine was dissolved directly in Tyrode’s solution. All experimental superfuses during recording were delivered by a home-built rapid solution exchange device [36].

2.4. Molecular modelling

Docking of ranolazine and lidocaine to hERG was conducted using homology models of the hERG pore encompassing the pore helix, selectivity filter and S6 helices. As with previous studies we found that an open channel model built onto the crystal structure of the bacterial K’ channel MthK (PDB: 1LNQ [37]) gave docking data broadly consistent with experimental mutagenesis data. This model is described elsewhere [34,38,39]. We have recently compared our MthK-based model with models based on putative C-type inactivated state structures of the KcSA bacterial channel and found that the MthK-based model accords more closely with experimental data on drug-binding for a range of structurally-diverse hERG blockers [39]. For other drugs that bind with moderate or weak affinity, binding modes obtained with a modified KvAP model (“Farid model”) [40] were similar to those described for the MthK model [34]. Here docking was performed with both the MthK and Farid models using the Flexdock module of SybylX2.0, which allows free sampling of protein side chain rotamers. Definition of the drug-binding pocket, selection of rotatable bonds, construction of starting configurations and choice of Genetic Algorithm parameters were performed as described previously [34,39]. Further information is available in the online supplementary information.

3. Results

3.1. Pharmacological sensitivity of WT hERG

The response of WT hERG to ranolazine was first determined using a ventricular action potential (AP) voltage command (AP clamp; Fig. 1Ai for representative traces and Fig. 1Aii for mean data). Application of 10 μM ranolazine produced IHERG suppression that was rapid in both onset and reversibility (Fig. 1Ai). Peak IHERG during the repolarizing phase of the AP was inhibited by 62.42 ± 2.03% (n = 6 cells). The rapid onset and washout of the effects of ranolazine are consistent with prior observations under conventional voltage clamp [26], in demonstrating that ranolazine inhibition both developed and washed out rapidly. We then used a conventional voltage step protocol, comprised of a 2-second depolarization to +20 mV followed by repolarization to −40 mV to elicit IHERG tails. Tail current (Itail) amplitude was measured relative to current elicited by a brief (50 ms) pre-pulse prior to the +20 mV test command [8,34,35,41]. Fig. 1B shows example records of IHERG in control and in the presence of 10 μM ranolazine (with 58% blockade of the IHERG tail in this example: (57.90 ± 1.29%, n = 6 cells; t-test, p > 0.05 compared to effects of this drug concentration under AP clamp)). The concentration–response relation for ranolazine inhibition of IHERG elicited by this protocol is shown in Fig. 1D (filled squares), yielding an IC₅₀ value of 8.03 ± 0.95 μM (h = 0.81 ± 0.07). Ranolazine thus produced hERG block at clinically relevant concentrations (2−6 μM [25]). For some drugs, reversal of the direction of K’ ion flux can reduce hERG block, due to electrostatic repulsion or ‘knock-off’ (e.g.[30,34,42,43]). Therefore, we also sought concentration–response data for ranolazine for inward hHERG currents at −120 mV, with a high (94 mM) extracellular K⁺ concentration ([K+]ₑ), using the protocol shown in Fig. 1C (lower panel). Representative current records
are shown in Fig. 1C (upper panel), with concentration–response data plotted in Fig. 1D (open circles). The sensitivity of inward \( I_{\text{hERG}} \) to ranolazine in 94 mM \([K^+]_e\) (an \( IC_{50} \) value of 8.47 ± 2.63 μM; \( h = 0.58 ± 0.10 \)) was similar to that for outward \( I_{\text{hERG}} \). Parallel experiments were performed on lidocaine (see online supplemental Fig. S2 and Table S2). The \( IC_{50} \) for outward \( I_{\text{hERG}} \) tail inhibition by lidocaine in 4 mM \([K^+]_e\) was 141.77 ± 9.81 μM (\( h = 0.98 ± 0.07 \)), with an IC\(_{50}\) for inward \( I_{\text{hERG}} \) in high \([K^+]_e\) increased to 735.32 ± 54.03 μM (\( h = 0.82 ± 0.05 \)). Thus lidocaine differed from ranolazine in its \( I_{\text{hERG}} \) blocking potency and sensitivity of inhibition to \([K^+]_e\) and to the direction of \( K^+ \) flux.

Further experiments were conducted under conventional voltage clamp in order to determine voltage dependence of WT \( I_{\text{hERG}} \) inhibition by ranolazine under our conditions (see supplemental Fig. S3 for details). Similar to a previous study [26], we observed ranolazine’s inhibitory action to show a degree of voltage dependence and to be accompanied by a modest left-ward shift in voltage-dependent activation of \( I_{\text{hERG}} \) (by ~4 mV at 10 μM; see supplementary Fig. S3).

### 3.2. The effect of inactivation attenuation on \( I_{\text{hERG}} \) inhibition

Intact hERG channel inactivation is important to the development of \( I_{\text{hERG}} \) inhibition by some (typically, though not exclusively, high affinity) drugs [8,30,44], although some agents with sub-μM and μM IC\(_{50}\) values do not depend critically on inactivation gating for inhibition to occur.
To our knowledge, no previous study has investigated the extent to which channel inactivation is required for \( I_{\text{hERG}} \) inhibition by ranolazine. S620T hERG channels lack inactivation and have been suggested to be useful for determination of true drug affinity for the open/activated channel [30]. Fig. 2A shows the effects of S620T on \( I_{\text{hERG}} \) block by ranolazine (Fig. 2Ai (representative traces) and Aii (concentration–response relations)). The drug's inhibitory potency was markedly reduced for S620T hERG compared to the WT channel, with an IC\(_{50}\) value of 582.65 ± 23.36 \( \mu \)M (\( h = 0.89 \pm 0.05 \)), ~73-fold the WT IC\(_{50}\). This mutation also had a marked effect on \( I_{\text{hERG}} \) block by lidocaine (see supplementary Fig. S4).

As ranolazine block of WT \( I_{\text{hERG}} \) occurs at clinically relevant concentrations, we further pursued the effect of inactivation attenuation by studying an additional, clinically relevant attenuated inactivation hERG mutant (N588K). Residue 588 lies in the S5-pore helix linker, which is distant from the hERG channel’s inner cavity. In contrast to the S620T mutation [30,44], the N588K mutation does not eliminate \( I_{\text{hERG}} \) inactivation completely, but results in positively shifted voltage-dependent inactivation of \( I_{\text{hERG}} \) by +60 to +90 mV [46,47]. This mutation is responsible for one form (SQT1) of the genetic short QT syndrome. It markedly attenuates the action of Class III agents such as sotalol, dofetilide and E-4031, but produces relatively small effects on inhibition by Class Ia drugs [30,48]. Fig. 2Bi shows representative N588K \( I_{\text{hERG}} \) traces in control and in 10 \( \mu \)M ranolazine. \( I_{\text{hERG}} \) block was attenuated compared to that of WT \( I_{\text{hERG}} \), but not as extensively as for the S620T mutation. Fig. 2Bii shows the concentration–response relation for inhibition of N588K \( I_{\text{hERG}} \) superimposed on that for WT \( I_{\text{hERG}} \). The derived IC\(_{50}\) value was 124.08 ± 7.39 \( \mu \)M (\( h = 0.81 \pm 0.05 \)) for N588K hERG, ~15-fold the corresponding value for WT \( I_{\text{hERG}} \). The reduction in ranolazine potency with this mutation is significantly greater than that reported for Class Ia or Class Ic drugs that have been studied previously [8,41,49,50].

3.3. Molecular determinants of ranolazine and lidocaine binding

The majority of hERG channel blockers studied bind within the channel’s inner cavity, interacting with one or both of two S6 aromatic residues (Y652, F656) [3]. There is some evidence that ranolazine
interacts with these aromatic residues [23]. We pursued this by measuring concentration–response data for Y652A and F656A hERG mutants with ranolazine and the structurally related lidocaine. Due to the known low expression and negatively shifted inactivation kinetics of F656A hERG [51], F656A IhERG was evaluated by measuring inward tail current in high [K+]e Tyrode (conditions comparable to those for WT IhERG in Fig. 1C). Figs. 3Ai and Aii show respectively the effects of ranolazine (10 μM) and lidocaine (300 μM) on F656A inward IhERG tails compared with WT IhERG controls. 10 μM ranolazine produced substantial inhibition of WT hERG tail current by 58.32 ± 3.87% (n = 6 cells). F656A hERG showed greatly attenuated IhERG inhibition, with fractional block of only 6.62 ± 0.62% by 10 μM ranolazine (n = 5 cells). Fig. 3Bi shows concentration–response relations for WT and F656A IhERG. The IC50 for F656A IhERG was 452.67 ± 12.07 μM (n = 5 cells per data-point.)

Fig. 3. Effect of S6 mutations on ranolazine and lidocaine block of IhERG. A. Expanded representative IhERG traces of WT hERG and F656A hERG in the absence (black line) and presence (grey line) of either 10 μM ranolazine (Ai) or 300 μM lidocaine (Aii). The current was evoked by the same protocol shown in Fig. 1C and was recorded in high K+ Tyrode’s solution (94 mM [K+]e). Horizontal arrows denote zero current. B. Concentration–response plots (mean ± SEM) for both WT (black line) and F656A (grey line) IhERG block by ranolazine (Bi) and lidocaine (Bii). Data were fitted with a Hill-equation. Error bars for some points are small and are obscured by the symbols. (n ≥ 5 cells per data-point.) C. Expanded example IhERG traces of WT hERG and Y652A hERG in the absence (black line) and presence (grey line) of either 10 μM ranolazine (Ci) or 300 μM lidocaine (Cii). The current was evoked by the same protocol shown in Fig. 1B and was recorded in normal Tyrode’s solution (4 mM [K+]e). Horizontal arrows denote zero current. D. Concentration–response plots (mean ± SEM) for both WT (black line) and Y652A (grey line) IhERG block by ranolazine (Di) and lidocaine (Dii). Data were fitted with a Hill-equation. Error bars for some points are small and are obscured by the symbols. (n ≥ 5 cells per data-point.)
0.68 ± 0.02), ~53-fold that of WT Iherged. Fig. 3Aii shows that 300 μM lidocaine inhibited F656A inward Iherged tails by 28.85 ± 1.80% (n = 8 cells) and WT Iherged tails by 30.19 ± 2.09% (n = 6 cells). Fig. 3Bii shows concentration–response data for lidocaine against F656A Iherged, which was closely superimposed over its WT control, with an IC50 value of 848.12 ± 65.54 μM (h = 0.85 ± 0.06), ~1.15-fold the value for WT. Thus, the F656 residue appeared to be a key binding determinant for ranolazine, but not for lidocaine.

The effect of Y652A on Iherged inhibition by ranolazine and lidocaine is shown in Figs. 3C and D (with the standard voltage protocol used for WT Iherged). Compared with the effect on WT Iherged (Fig. 3Ci), inhibition of Y652A Iherged by 10 μM ranolazine was significantly reduced (Itail inhibition of ~19%). The concentration-dependence of mean fractional block is plotted in Fig. 3Di. The IC50 for inhibition of Y652A Iherged by ranolazine was 173.62 ± 39.73 μM (h = 0.64 ± 0.14), ~22-fold the corresponding value for WT hERG. Figs. 3Cii and Dii show the effect of 300 μM lidocaine, with a reduction of the Y652A Itail by ~36.64% compared to ~70.11% for the WT Itail. The IC50 for inhibition of Y652A Iherged by lidocaine was 554.33 ± 34.03 μM (h = 0.95 ± 0.06), ~3.8-fold the corresponding value for WT hERG. Thus, the Y652A mutation had a marked effect on ranolazine block of Iherged, but only a modest effect on the action of lidocaine.

Amino acids located at the base of the pore helix (T623, S624 and V625) have been identified as important components of drug-binding sites for some [51–53] though not all [34,38] hERG blockers. To investigate the roles of these residues in ranolazine and lidocaine block the T623A, S624A and V625A mutations were studied. T623A Iherged was studied using the same voltage protocol and conditions as used for

Fig. 4. Effect of pore helix mutations on ranolazine and lidocaine block of Iherged. A. Fitted concentration–response relations for WT hERG (black line) and T623A hERG (grey line) inhibition by ranolazine (Ai) and lidocaine (Aii). Panels B, and C show equivalent data for T624A, and V625A, respectively. Error bars for some points are small and are obscured by the symbols. For all plots n ≥ 5 cells per data-point.
F656A I_{hERG}. Figs. 4Ai and Aii respectively show the concentration–response relations for ranolazine and lidocaine against WT and T623A I_{hERG}. The IC_{50} for T623A I_{hERG} block by ranolazine was 159.21 ± 2.42 μM (h = 0.52 ± 0.006), ~19-fold its WT control (Fig. 4A), whilst that for lidocaine was 746.48 ± 70.40 μM (h = 0.81 ± 0.07), 1.01-fold its WT control (Fig. 4Aii). The effect of S624A hERG on I_{hERG} block by ranolazine was investigated using the standard voltage protocol (shown in Fig. 1B). Figs. 4Bi and Bii show the concentration–response relations respectively for ranolazine and lidocaine for S624A I_{hERG}. The IC_{50} for S624A I_{hERG} block by ranolazine was 61.53 ± 12.93 μM (h = 0.93 ± 0.19), ~8-fold that of WT I_{hERG} (Fig. 4Bi), whilst that for lidocaine was 107.59 ± 14.18 μM (h = 1.03 ± 0.14), ~0.76-fold that of WT I_{hERG}. V625 is located within the K⁺ signature sequence of hERG (SVGFG) and mutation to alanine reduces the selectivity of hERG channel for K⁺ [51]. This mutant requires to be studied by measuring the inward hERG tail current at −120 mV. Fig. 4Ci shows concentration-response relations for WT and V625A I_{hERG} inhibition by ranolazine, with an IC_{50} of 63.14 ± 6.88 μM (h = 0.58 ± 0.04), ~8-fold that of WT I_{hERG}. Fig. 4Cii shows similar data for lidocaine, which for V625A I_{hERG} had an IC_{50} of 1580 ± 75.48 μM (h = 0.51 ± 0.01), 2.68-fold its WT control. Thus, each of the pore helix mutants had modest effects on ranolazine block of I_{hERG}, whilst for lidocaine T623A and S624A were essentially without effect and V625A had only a small effect.

The effects of all of the studied hERG mutants on the potency of inhibition of I_{hERG} by both ranolazine and lidocaine are summarized in supplementary Tables S1 and S2.

3.4. Molecular modelling of ranolazine and lidocaine action

Docking of lidocaine and ranolazine into the MthK and Farid models produced similar low energy score docked states. In both models the positively-charged tertiary amine common to both blockers is localized near the binding site for a K⁺ ion observed in crystal structures of potassium channels (Fig. 5). K⁺ ions are stabilised here by the focused...
electrostatic field arising from the C-terminal helix dipoles of the pore helices. In this configuration each blocker can make additional interactions with aromatic side chains of Y652 and F656. Neither drug made direct contact with V625 or S620 (Fig. 5), consistent with previous evidence that effects of S620T and V625A mutations on drug-binding result from indirect effects on hERG pore conformations [e.g. [54]]. Although both drugs lie high in the pore cavity just below the selectivity filter, the orientations of the amide group differed, with the ranolazine amide lying horizontally in a position where it may hydrogen bond with the S624 side chain (Figs. 5B, E, F). The lidocaine amide was more vertically oriented and unable to hydrogen bond with the S624 OH group (Figs. 5A, C, D). These observations are consistent with the mutagenesis data for S624A (Figs. 4Bi, Bii). Similarly, lidocaine made no interactions with T623 (Figs. 5A, C, D) consistent with a lack of effect of T623A on lidocaine block (Fig. 4Aii). In some poses (e.g. Fig. 5E) ranolazine approached close to a T623 side chain; this may be relevant to the reduced block of T623A (Fig. 4A).

The structural basis for enhanced drug-binding to the hERG inactivated state has been attributed to the conformation of the S6 helix in which side chains, particularly Y652 and F656, are optimally positioned in the inactivated state to interact with drug molecules in the pore cavity [55]. The large suppression of ranolazine block in hERG S620T (Fig. 2Bi) is consistent with the multiple interactions of ranolazine moieties with Y652 and F656 side chains in WT hERG (Figs. 5E, F), and the attenuation of these interactions by non-optional arrangements of aromatic side chains in S620T; this accounts for the observation that S620T produces a larger attenuation of block than either Y652A or F656A alone.

Ranolazine inhibition of I_{hERG} was substantially suppressed by the S620T mutation (supplemental Fig. S4). However, suppression of I_{hERG} block was negligible for F656A hERG (Fig. 3Bii) and only ~4-fold in Y652A (Fig. 3Dii), indicating that neither of these side chains is essential for block. Docking of lidocaine into hERG models carrying either the Y652A or F656A mutation showed that these observations may be understood in terms of a degeneracy in binding partners for the lidocaine aromatic group (supplemental Fig. S5); the lidocaine aromatic ring can make interactions with either a Y652 (in the case of F656A) or F656 (in Y652A) aromatic side chain. However if optimal configurations of both Y652 and F656 side chains are lost in S620T hERG, low affinity lidocaine block is considerably suppressed. By contrast, docking of ranolazine to models carrying either the Y652A or F656A mutation demonstrated that interactions with both Y652 and F656 aromatic side chains are required for optimal drug–pore interactions to occur (supplemental Fig. S6).

4. Discussion

4.1. Comparison with previous studies

The potency of I_{hERG} block by ranolazine reported here (IC$_{50}$ of ~8 μM) is similar to that reported previously by Rajamani and colleagues (~12 μM) [26]. We found that ranolazine was nearly 18-fold more potent than lidocaine at inhibiting I_{hERG}. Consistent with only a weak propensity of lidocaine to inhibit I_{hERG} [11], ranolazine has previously been suggested to compete with the methansulphonanilide E-4031 for a binding site on hERG [23], with some data supporting involvement of S6 aromatic residues in this action [23]. However, to our knowledge, no prior study has provided information on the role of inactivation gating in ranolazine’s inhibitory action, nor a detailed analysis of structural determinants of inhibition. Nor, in contrast to its interactions with Na$^+$ channels [31,56], has any prior comparative analysis in respect of hERG block been conducted with lidocaine.

4.2. Role of inactivation in I_{hERG} block by ranolazine

The potency of ranolazine’s inhibition of I_{hERG} reported here is similar to that of the Class 1a antiarrhythmic drug disopyramide (~7–11 μM [8,34]), but ranolazine differs markedly from disopyramide in two key respects: (i) disopyramide block is largely insensitive to mutations near the bottom of the selectivity filter (T623, S624, V625); (ii) disopyramide shows only a weak dependence of block on intact inactivation [8,34]. These characteristics are shared by other Class Ia/c drugs [8,38]. By contrast, the S620T–inactivation deficient channel showed markedly reduced sensitivity to ranolazine (and lidocaine). The location of the S620 residue on the pore helix and oriented towards the extracellular surface of the pore (Figs. 5A, B) excludes a direct interaction of this residue with drugs in the pore. Instead, the effect of S620T likely arises from the loss of configurations of side chains (especially Y652 and F656) upon inactivation gating that are favourable for drug interactions [55]. Indeed, titration of inactivation loss appears to produce a graded reduction in blocking potency of some drugs [8]. For ranolazine, the 72-fold reduction in the block of the S620T hERG mutant is similar to the sum of the individual reductions in block of hERG Y652A (22-fold) and F656A (53-fold), consistent with the expectation that ranolazine requires interactions with aromatic side chains of both residues to express full block. This is consistent with docking which shows simultaneous aromatic interactions with both Y652 and F656 residues (Fig. 5). And, the loss of interactions in each of Y652A and F656A (Fig. 5). In contrast with the N588D (neutral → negative charge substitution) long QT syndrome [56] mutation in the external S5–Pore linker, which results in a negative-shift in voltage-dependent I_{hERG} inactivation [57], the N588K (neutral → positive charge substitution) short QT syndrome mutant produces a marked positive shift in voltage-dependent inactivation, but a change that is less extensive than for S620T [8,30]. Less extensive reduction in ranolazine block for N588K likely reflects the fact that inactivation is incompletely removed for N588K–hERG [8], so that interactions with aromatic residues are less extensively perturbed than for S620T. Moreover, in additional experiments performed on WT hERG, in which ranolazine was washed off gated hERG channels during sustained membrane depolarization (data not shown), the rate of current recovery was slowed at more depolarized voltages at which inactivation was promoted. This supports a role for inactivation gating in stabilizing ranolazine binding within the channel pore. In a previous investigation of the comparative pharmacology of the N588K short QT hERG mutant, there was a correlation between hERG blocking potency and sensitivity of block to attenuated inactivation [8]. However, despite a similar I_{hERG} blocking potency to disopyramide [8,34], ranolazine shows a much greater sensitivity to inactivation attenuation (~15-fold WT IC$_{50}$ for N588K) than disopyramide (~1.5 WT IC$_{50}$ for N588K) [8]. This argues against a fixed relationship between WT I_{hERG} blocking potency and dependence of blockade on inactivation. Our docking simulations (Fig. 5) indicated that ranolazine cannot interact directly with V625 whilst interacting with S6 aromatic residues. Consistent with suggestions for other drugs (e.g. [54]), the modest effect of V652A on ranolazine block of hERG is likely to arise from indirect effects of this mutation on the positioning of drug-binding residues.

4.3. Comparison between ranolazine and lidocaine

Lidocaine shares structural features with ranolazine but is a smaller molecule (see supplementary Fig. S1). Our findings indicate that the two drugs differ in their actions on hERG in the following respects: (i) inhibitory potency (ranolazine > lidocaine); (ii) sensitivity of inhibition to pore helix and S6 helix mutations; (iii) sensitivity of inhibition to potassium flux.

Mutagenesis and docking simulations demonstrate the potential for direct interactions between ranolazine and T623 and S624 residues, with hydrogen-bonding possible with S624 side chains. These interactions are absent for lidocaine. Ranolazine with two aromatic rings can make multiple interactions with Y652 and F656 side chains whereas the smaller lidocaine molecule makes fewer interactions (supplementary Fig. S7). Collectively, these factors can account for the greater inhibitory
potency of ranolazine than of lidocaine. As discussed above, the strong dependence of \(i_{hERG}\) inhibition by ranolazine on intact inactivation kinetics can be accounted for by optimal positioning of \(S(6)\) binding residues. For lidocaine, the effect of the S620T mutation (a 48-fold higher IC\(_{50}\) in potency) was much greater than that of either Y652A (3.84-fold WT IC\(_{50}\)) or F656A (1.15-fold WT IC\(_{50}\)) or their sum. Lidocaine possesses both an aromatic ring and a charged aliphatic amine group that can be anticipated to interact with aromatic side chains (through \(\pi-\pi\) and cation–\(\pi\) interactions respectively). Indeed, block of voltage gated Na\(^+\) channels by local anesthetics relies on Phe and Tyr residues (equivalent to F1670 and Y1767 in Nav1.5 isoform-1) in the S6 segment of domain 4. [58,59] with the Phe likely interacting directly with lidocaine through a cation–\(\pi\) interaction [60]. Ranolazine’s inhibitory action on both peak and persistent Na\(^+\) current is sensitive to mutation of F1760 [20], indicating a common binding site for lidocaine and ranolazine on Na\(_{\text{v}}\) channels (although the underlying molecular interactions may differ [58]). Thus, it seems reasonable to propose that lidocaine is also likely to interact with S6 aromatic residues of hERG and our docking data (Figs. 5 and S5) support such a conclusion. Mutagenesis and docking data can be reconciled by degeneracy of lidocaine interactions with the two aromatic residues; the docking simulations suggest that the aromatic ring of lidocaine can interact with either F656 or Y652 residues. We attribute the marked reduction in lidocaine’s \(i_{hERG}\) inhibitory action by the S620T inactivation mutant to the loss of favourable configurations of both Y652 and F656 in S620T, with the consequence that the aromatic group of lidocaine is unable to make any aromatic interactions within the pore cavity (Figs. 5 and S5).

Consistent with our interpretations for hERG (Figs. 5 and S5), modeling of lidocaine binding to the open state Na\(_{\text{v}}\) channel supports binding poses in which the positively-charged amino group is located below the selectivity filter where the pore helix negative dipole charges are focussed [60]. This binding mode makes lidocaine block susceptible to competing charge effects (in the case of Na\(_{\text{v}}\) as a result of charge repulsion between lidocaine and a Na\(^+\) ion in its DEKA binding site in non-inactivated states [60]). This is manifest in hERG as a reduction of lidocaine block by competition from K\(^+\) ions under conditions of inward ion flux (Figs. 1Cii and Dii). The docked poses for lidocaine (Figs. 5 and S5) suggest an explanation for the reduction in hERG block by inward K\(^+\) flux, since binding interactions comprise only the tertiary amino group with the pore helix dipole charges and the aromatic ring with either a Y652 or F656 side chain. The electrostatic interaction constitutes a large proportion of the total contributions to binding so that competition with permeant ions for the K\(^+\) binding site significantly reduces lidocaine affinity. The electrostatic component makes a much smaller contribution to binding for ranolazine which is relatively resistant to the effects of permeant ions.

### 4.4. Significance for clinical drug use and development

Recent human subject data indicate that plasma concentrations (\(C_{\text{max}}\)) of ranolazine of 741.5 ng/ml and 2328.7 ng/ml (~1.73 and ~4.05 \(\mu\)M) are attained after single oral doses of 500 mg and 1500 mg respectively [61]. These levels are in good agreement with a clinical plasma concentration range of 2–6 \(\mu\)M quoted by Rajamani and colleagues [25]. Thus, partial \(i_{hERG}/i_{K_0}\) inhibition can be expected to occur during clinical use of ranolazine. By contrast, plasma lidocaine levels during clinical use approximate ~7–15 \(\mu\)M [62,63] and, with an IC\(_{50}\) for \(i_{hERG}\) inhibition close to 142 \(\mu\)M, lidocaine is anticipated to produce little or no \(i_{hERG}/i_{K_0}\) inhibition during therapeutic use. Our results explain why ranolazine is able to inhibit \(hERG/i_{K_0}\) at clinically relevant concentrations, whilst structurally similar lidocaine only produces significant inhibition at concentrations markedly exceeding the normal clinical range. Although ranolazine can exert weak actions on other ionic currents, the dominant ventricular ion channel effects of ranolazine at therapeutic concentrations are inhibition of \(i_{hLA,T}\) and \(i_{K_0}\), whilst comparatively atrioselective inhibition of rapid, peak \(i_{Na}\) [24] additionally contributes importantly to the atrial actions of the drug [64,65]. The \(i_{K_0}\) blocking effect of ranolazine has been proposed to be significant to the drug’s action against AF [24,28,64]. The greater atrial than ventricular APD-prolonging effect of the drug may result from differences in AP configuration (and roles of \(hLA_{\text{late}}\)) in the two tissue types, whilst the drug’s \(i_{K_0}\) blockade enhances inactivated state Na\(^+\) channel block and therefore ranolazine’s overall effectiveness [24,64].

The larger size of ranolazine gives it potential for a greater range of interactions with hERG pore side chains compared to lidocaine, and enables interaction of its two aromatic groups with side chains of both Y652 and F656. Our analysis indicates that low potency lidocaine block has low susceptibility to either of the Y652A and F656A mutations because either aromatic side chain can “compensate” for loss of the other. Given that combining Na\(^+\) channel and \(i_{K_0}\) actions may hold promise for antiarrhythmic drug rate-dependency [27,66–68], the comparative information on ranolazine and lidocaine in this study in respect of hERG block may feasibly have utility for future design of agents combining \(i_{hERG}\) and \(i_{K_0}\) inhibition, through modification(s) of the lidocaine template that is common to both drugs. Our data in respect of ranolazine are also of significance to pharmacological treatment of the hERG-linked (SQT1) variant of the genetic short QT syndrome. Patients with N588K-hERG-linked SQT1 are at risk of ventricular arrhythmia and sudden death and are usually treated an implantable cardioverter defibrillator, which itself carries a risk of inappropriate shocks [69]. This makes antiarrhythmic treatment an attractive adjunct approach. However, ranolazine inhibition of \(i_{hERG}\) at clinically relevant drug concentrations appears to require intact inactivation and, in particular, the reduction in ranolazine potency with the N588K hERG mutation seen here is significantly greater than that reported for Class 1a or Class 1c drugs that have been studied previously [8,41,49,50]. Consequently, ranolazine is unlikely to be a suitable alternative to Class 1a antiarrhythmic agents currently used for the pharmacological treatment of SQT1 [70].

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### Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.yjmcc.2014.05.013.

Conflict of interest statement

None.

### References


