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Probing the bradycardic drug binding receptor of HCN-encoded pacemaker channels

Yau-Chi Chan · Kai Wang · Ka Wing Au · Chu-Pak Lau · Hung-Fat Tse · Ronald A. Li

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Abstract I_f (or I_h), encoded by the hyperpolarization-activated, cyclic nucleotide-gated (HCN1–4) channel gene family, contributes significantly to cardiac pacing. Bradycardic agents such as ZD7288 that target HCN channels have been developed, but the molecular configuration of their receptor is poorly defined. Here, we probed the drug receptor by systematically introducing alanine scanning substitutions into the selectivity filter (C347A, I348A, G349A, Y350A, G351A in the P-loop), outer (P355A, V356A, S357A, M358A in the P-S6 linker), and inner (M377A, F378A, V379A in S6) pore vestibules of HCN1 channels. When heterologously expressed in human embryonic kidney 293 cells for patch-clamp recordings, I348A, G349A, Y350A, G351A, P355A, and V356A did not produce measurable currents. The half-blocking concentration (IC_{50}) of wild type (WT) for ZD7288 was 25.8±9.7 μM. While the IC_{50} of M358A was identical to WT, those of C347A, S357A, F378A, and V379A markedly increased to 137.6±56.4, 113.3±34.1, 587.1±167.5, and 1726.3±673.4 μM, respectively (p<0.05). Despite the proximity of the S6 residues studied, M377A was hypersensitive (IC_{50}=5.1±0.7 μM; p<0.05) implicating site specificity. To explore the energetic interactions among the S6 residues, double and triple substitutions (M377A/F378A, M377A/V379A, F378A/V379A, and M377A/F378A/V379A) were generated for thermodynamic cycle analysis. Specific interactions with coupling energies (ΔΔG) >1 kT for M377–F378 and F378–V379 but not M377–V379 were identified. Based on these new data and others, we proposed a refined drug-blocking model that may lead to improved antiarrhythmics and bioartificial pacemaker designs.

Keywords Drug · Inner pore · ZD7288 · Pacemaker channels · HCN

Introduction

Pacemaker activity, the generation of spontaneous cellular electrical rhythms, governs numerous biological processes from the autonomous beating of the heart to respiratory rhythms and sleep cycles [31]. In the heart, abnormal pacing leads to various forms of arrhythmias (e.g., sick sinus syndrome). I_f (or I_h in noncardiac tissues), encoded by the hyperpolarization-activated, cyclic nucleotide-gated (HCN1–4) channel gene family, is a key depolarizing current in cardiac pacing that is activated by hyperpolarization (rather than depolarization of classical voltage-gated...
ion channels) [15, 23, 27]. Familial mutations of the human HCN genes have been described [24].

Structurally, each HCN channel consists of four homologous monomers pseudosymmetrically arranged around a central pore, resembling voltage-gated K+ (Kv) channels; each of the four internal repeats is made up of six transmembrane segments (S1–S6; Fig. 1). The region between S5 and S6, or the so-called P-loop, inserts back into the membrane to form part of the pore. Indeed, the pore is analogous to the active site of an enzyme where major functional (e.g., ionic selectivity, conductance and gating) and pharmacological determinants are located [31].

Given the physiological importance of If, bradycardic drugs such as ZD7288 and ivabradine that target HCN channels have been developed. However, the molecular constituents of the HCN channel drug binding receptor have not been fully elucidated. A better understanding is crucial for designing more effective, HCN-specific drugs. For voltage-gated K+ and Na+ channels, their drug binding sites have been identified to reside in the pore regions [18]. Given the structural similarities between HCN and Kv channels, it has been hypothesized that the drug receptor of HCN channels is similarly located in the pore [5, 6, 12]. Shin and colleagues demonstrate that ZD7288 applied from the cytoplasmic side can enter and leave the inner pore of HCN1 channels only at voltages where the activation gate is opened [30]. Using site-directed mutagenesis, Sanguinetti and colleagues report that the S6 residues A425 and I432 of HCN2 channels (the analogous residues are A372 and V379 in HCN1) are primary determinants for ZD7288 block as well as their implications in bradycardic drug and bioartificial pacemaker designs.

Materials and methods

Molecular biology and heterologous expression Murine HCN1 cDNA was cloned from mouse brain RNA [27] and subcloned into the mammalian expression vector pCI (Promega, Madison, WI, USA). Desired substitutions were constructed as previously described [38]. HCN1 channels were transiently expressed in human embryonic kidney 293 cells (HEK293) using Lipofectamine Plus 2000 (Invitrogen, Carlsbad, CA, USA). For identifying HCN1-expressing cells, the channel plasmid was cotransfected with pCI-green fluorescent protein (GFP) with a ratio of 10:1. Twenty-four hours to 48 h after transfection, cells were trypsinized and plated on glass-bottomed culture dishes 2 h prior to patch-clamp or immunohistochemical experiments.

Electrophysiology Whole-cell patch-clamp recordings were performed using an EPC-10 patch-clamp amplifier (HEKA electronic, Heidelberg, Germany). Only GFP-expressing HEK293 cells as identified by their epifluorescence with an excitation wavelength of 488 nm were selected for experiments. All recordings were performed at room temperature in a bath solution containing (in mM): 110 NaCl, 30 KCl, 1.8 CaCl2, 0.5 MgCl2, 5 HEPES, and 10 glucose, pH adjusted to 7.4 with NaOH. The internal solution contained (in mM): 10 NaCl, 130 KCl, 0.5 MgCl2, 5 HEPES, 1
a: -140 mV  
b: -100 mV  
c: -70 mV

P-loop
C347A  
V348A

P-S6
P355A  
V356A

S6
M377A  
V379A

G349A
S357A  
V379I

Y350A
M358A  
V379I

G351A
EGTA, and 5 MgATP, pH adjusted to 7.3 with KOH. The electrode tip and series resistances were, respectively, ~3–4 and <10 MΩ. The sampling frequency was 1.25 kHz. Cell capacitance and series resistance compensation were not used. The cycle length of the protocols was 4.6 s. The potentials were not corrected for the liquid junction potentials of +20.3 mV.

**Experimental protocols and data analysis** The steady-state current–voltage (I–V) relationship was determined by plotting currents measured at the end of a 3-s pulse ranging from −140 to −30 mV from a holding potential of −30 mV against the test potentials. The voltage dependence of HCN channel activation was assessed by plotting tail currents measured immediately after pulsing to −140 mV as a function of the preceding 3-s test pulse normalized to the maximum tail current recorded. The data were fitted to the Boltzmann functions using the Marquardt–Levenberg algorithm in a nonlinear least squares procedure:

\[
m = \frac{1}{1 + \exp\left(\frac{V_t - V_{1/2}}{k}\right)}
\]

where \(V_t\) is the test potential; \(V_{1/2}\) is the half-point of the relationship; \(k=RT/\Delta F\) is the slope factor, and \(R, T, \Delta, \text{and } F\) have their usual meanings.

For half-blocking concentrations (IC50), the following binding isotherm was used:

\[
\frac{I}{I_0} = 1/(1 + ([\text{blocker}]/\text{IC50})^n)
\]

**Fig. 2 (continued)**

**Fig. 3** Steady-state current voltage and activation curves. a Effects of Ala and Ile substitutions on HCN1 steady-state current–voltage relationships. b Representative tracings and steady-state activation relationships
where \( n \) is the Hill coefficient; \( I_0 \) and \( I \) are the peak currents measured at the voltage indicated before and after blocker application, respectively. All IC\(_{50}\) values reported were calculated from individual determinations.

For thermodynamic cycle analysis [17], coupling coefficients (\( \Omega \)) and interaction energies (\( \Delta G \)) for various mutant channel pairs were calculated from the corresponding IC\(_{50}\) values with the equations below:

\[
\Omega = \left( \frac{IC^{\text{ConstructA}}_{50}}{IC^{\text{ConstructAB}}_{50}} \right) \left( \frac{IC^{\text{WT}}_{50}}{IC^{\text{ConstructB}}_{50}} \right)
\]

\[
\Delta \Delta G = RT \ln \Omega
\]

where \( R \) is the gas constant \((k \times N_A)\), and \( T \) is the absolute temperature. The standard errors for \( \Delta \Delta G \) were estimated by dividing the square root of the sum of the variances of the \( RT \ln IC_{50} \) means by the square root of the degree of freedom. Significant interactions were defined as \( \Delta \Delta G > 1 \text{ kcal/mol} \) as we and others previously reported [20].

All data presented were the means±SEM. Statistical significance (\( p<0.05 \)) was determined using an unpaired Student’s \( t \) test.

**Immunostaining** HEK293 cells were fixed in 4% paraformaldehyde for 15 min at 21°C, washed with phosphate-buffered saline (PBS), and permeabilized with 0.1% Triton-X-100/PBS. The cells were then blocked with 10% bovine serum albumin with 4% goat serum in PBS for 2 h at 21°C. Fixed cells were incubated with the HCN1 rabbit polyclonal antibodies (Alomone Labs, Jerusalem, Israel) at a dilution of 1:200 overnight at 4°C, followed by incubation with fluorescent-labeled secondary antibodies for 50 min at 21°C and visualization by fluorescent microscopy.

**Results**

**Effects of alanine (Ala) substitutions on HCN1** As a first step, we generated a total of 12 singly Ala-substituted HCN1 pore constructs that correspond to these regions: C347A, I348A, G349A, Y350A, and G351A in the ascending limb of the P-loop, P355A, V356A, S357A, and M358A in the P-S6 linker that forms part of the extrapore and M377A, F378A, and V379A in the S6 segment that constitutes part of the cytoplasmic pore mouth (Fig. 2a). Of note, I432 of HCN2, the analogous residue to I348 of HCN1, has been previously reported to involve in IC\(_{50}\) values with the equations below:

\[
\Delta \Delta IC = \frac{IC_{50A} - IC_{50B}}{IC_{50W}}
\]

\[
\Delta \Delta I = \frac{I_{50A} - I_{50B}}{I_{50W}}
\]

where

\[
\Delta \Delta G = \frac{R \ln \Omega}{RT}
\]

and

\[
\Delta \Delta I = \frac{R \ln \Omega}{RT}
\]

were calculated from individual determinations. The corresponding steady-state current–voltage (\( I-V \)) relationships are shown in Fig. 3a. The different activation thresholds hint that the gating properties of some of these Ala-substituted pore constructs have been altered. To test this notion, we next examined their steady-state activation properties (Fig. 3b). Indeed, all of C347A (P-loop), S357A, and M358A (P-S6), M377A, and V379A (S6) similarly and significantly hyperpolarized the steady-state activation midpoint (\( V_{1/2, \text{con}=} \) of WT from −62.6±1.4 mV (\( n=6 \)) to −82.1±4.7 mV (\( n=3 \)), −76.1±0.3 mV (\( n=3 \)), −70.8±1.6 mV (\( n=5 \)), −77.4±1.6 mV (\( n=4 \)) and −97.8±2.6 mV (\( n=8 \)) mV, respectively (\( p<0.05 \)). By contrast, F378A (of S6) caused a noticeable opposite depolarizing shift (\( V_{1/2, \text{con}=} \)−56.2±3.7 mV, \( n=3 \)) although the difference did not reach statistical significance (\( p=0.09 \)). Of note, C347A, F378A, and V379A increased the slope factor (\( k \)) by ~2-fold, twofold, and fivefold, respectively (\( p<0.05 \)).

**Effects of Ala substitutions on current blockade by ZD7288** To investigate the effect of our pore substitutions on drug block, we started by characterizing the response of WT HCN1 to ZD7288. Figure 4a shows that ZD7288 dose-dependently reduced the maximum steady-state current of WT at −140 mV. The half-blocking concentration or IC\(_{50}\) estimated from the binding curve was 25.8±9.7 \( \mu \)M (\( n=5 \)). The steady-state \( I-V \) and activation curves recorded in the absence and presence of ZD7288 are given in Fig. 4b, c, respectively. Figure 5 summarizes the effect of Ala substitutions on ZD7288 blockade. C347A, S357A, F378A, and V379A significantly shifted the binding curve in the rightward direction. Their IC\(_{50}\) were 137.6±56.4 (\( n=4 \), 113.3±34.1 (\( n=5 \), 587.1±167.5 (\( n=3 \)), and 1,726.3±673.4 \( \mu \)M (\( n=3 \)), respectively. Despite the proximity of M377 to F378 and V379 in the primary sequence of S6, its Ala substitution exerted an opposite effect on ZD7288 block.
Control
ZD7288 30 µM
ZD7288 100 µM

Normalized Current
Vm (mV)

IC50 = 25.8 ± 9.7 µM

[ZD7288] (µM)

M377A Control
ZD7288 3 µM
ZD7288 1 mM


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Fig. 5  Effects of Ala substitutions on ZD7288 block.  

**a**  
Upper panel: current tracings recorded at $-140$ mV in the presence of ZD7288 at different concentrations.  
Lower panel: the dose–response curves.

**b**  
Summary of IC$_{50}$.  
Dashed line indicates the level of WT sensitivity (*, $p<0.05$)


Energetic interactions among the MFV<sub>377–379</sub> triplet in S6

Among all the pore sites investigated, the S6 residues of the MFV<sub>377–379</sub> triplet, which are in close proximity to each other, when Ala-substituted produced the most significant changes in their responses to ZD7288 and gating properties, consistent with the role of S6 residues in drug binding. To explore their energetic interactions, three double (M377A/F378A, M377A/V379A, and F378A/V379A) and triple substitutions (M377A/F378A/V379A) were generated for thermodynamic cycle analysis. All constructs led to robust current expression. As for ZD7288 blockade assessed at −140 mV, the IC<sub>50</sub> values as estimated from the dose–response curves of M377A/F378A, M377A/V379A, F378A/V379A, and M377A/F378A/V379A channels were 2,253.8±378.6 (n=6), 313.5±79.4 (n=4), 1,267.0±484.1 (n=6), and 2,027.4±1,100.9 μM (n=4), respectively (Fig. 6; p<0.05). To calculate their interaction energies with each other, a representative cycle analysis was given in Fig. 7a as illustration. The coupling coefficient (Ω) for the interaction pair M377–F378 calculated from WT, M377A, F378A, and M377A/F378A was 0.05±0.01. Using the corresponding single and double substitutions, the M377–V379 and F378–V379 pairs were calculated to have Ωs of 1.09±0.21 and 31.0±9.09, respectively. Their absolute interacting energies were −1.77±0.09, 0.05±0.13, and 2.03±0.18 kcal/mol, respectively (Fig. 7b). These results indicate that significant interactions exist for the pairs M377–F378 and F378–V379 but not M377–V379.

As a means to verify the specificity of our results presented above, we constructed the triple construct M377A/F378A/V379A for further cycle analyses. To calculate the interaction of the pair M377–F378 in the background of V379A, the combination of V379A, M377A/V379A, F378A/V379A, and M377A/F378A/V379A was employed. The Ω and ΔΔG calculated for the pair were 0.11±0.01 and −1.31±0.10 kcal/mol (or −2.21±0.15 kcal/mol), respectively. Using the same approach, the Ω and ΔΔG for the pairs of F378–V379 in the M377A background and M377–V379 in the F378A background were 68.17±9.09 and 4.22±0.54 and 2.40±0.21 and 0.88±0.24 kcal/mol, respectively. In other words, the same rank order for interactions as those calculated from WT and single and double substitutions was observed.

V379I displayed WT gating and ZD7288 phenotypes

Isoform-specific differences in bradycardic drug block of HCN1 and HCN4 by the ivabradine have been reported [7]. Of note, the analogous residues of V379 of HCN1 are isoleucines in HCN2, HCN3, and HCN4 (Fig. 1). However, V379I displayed gating and ZD7288 sensitivity not different from WT (Figs. 2, 3, and 5).

Discussion

In the present study, we probed the molecular constituents of the drug binding receptor in HCN channels by Ala scanning mutagenesis of three distinct pore regions: the selectivity filter and the outer and the inner pore vestibules. Among the pore residues examined, the S6 triplet MFV<sub>377–379</sub> appears to play the most significant role in determining the WT drug-blocking phenotype. When Ala-substituted, residues M377, F378, and V379 significantly affected the sensitivity of HCN1 channels to the bradycardic drug ZD7288 by up to 67-fold (versus the −3-fold differences of the P- and P-S6 constructs). This finding is consistent with the notion that the inner pore region is crucial for drug binding as demonstrated by numerous previous Kv and Na<sup>+</sup> channel studies [18]. Despite their proximity in the primary sequence, M377A enhanced whereas F378A and V379A reduced ZD7288 block. Furthermore, thermodynamic cycle analysis revealed that the pairs M377–F378 and F378–V379, but not M377–V379, energetically interact. Importantly, whereas M377 and F378 negatively interact with other in drug binding, the interaction between residues F378 and V379 is positive. The same common pattern was observed when channel backgrounds other than WT were employed for cycle analyses. These differential effects strongly support that the observed changes in ZD7288 block were highly site specific.

Sequence alignments and comparison with the known crystal structures of K<sub>v</sub> channels suggest that the side chain of S6–V379 of HCN1 faces the aqueous phase of the cytoplasmic pore. In Shaker K<sup>+</sup> channels, the hydrophobic binding pocket for quaternary ammonium compounds consists of T469 in S6 [13]; the analogous residue of Shaker’s T469 in HCN1 is C374. Our results indicate that C347A display a fourfold reduction in ZD7288 sensitivity. Taken together with the present findings that the S6 substitutions F378A and V379A (but not M377A) reduce ZD7288 block, we propose that residues F378 and V379 are located on the same side of the S6-MFV<sub>377–379</sub> helical...
turn that is exposed to the cytoplasmic pore. This spatial arrangement is consistent with our thermodynamic cycle analysis that energetic interactions exist only for the pairs M377–F378 and F378–V379 but not M377–V379. Along with the P-loop residue C374, the F378/V379 helical face forms a receptor pocket to which ZD7288 binds. Although the S6 residue M377 faces a different side of the S6-MFV377–379 helix, it regulates the pathway by which the drug molecule travels to its F378–V379 binding pocket: when M377 is substituted by the smaller alanine, drug access is improved, sterically and/or by relieving some negative interactions, thereby facilitating ZD7288 block. This proposed mechanism is also consistent with the negative coupling identified for the M377–F378 pair. Our model is schematically presented in Fig. 8.

![Fig. 6 Effects of double and triple Ala substitutions on ZD7288 block.](image)

**Upper panel:** the current tracings recorded at −140 mV in the presence of ZD7288 at different concentrations. **Middle panel:** the dose–response curves for ZD7288 block of the WT, double and triple HCN1 channels. **Lower panel:** bar graph summarizing the IC₅₀. Dashed line indicates the level of WT sensitivity. Asterisks indicate statistical differences (*)p<0.05) compared with WT.
HCN2, a residue homologous to Val379 in the MFV_{377–379} triplet of HCN1, has also been shown to be involved in cilobradine binding [12]. Further investigations will be needed to test whether there is a single common receptor in HCN channels for these drugs or there are multiple overlapping and/or nonoverlapping drug binding sites. The interacting MFV triplet that we have identified in the present study appears to be a prime candidate for testing these possibilities.

Though we have identified key residues that determine ZD7288 block of HCN1 channels and energetically quantified their interactions, the nature of their side chain interactions with ZD7288 remains to be explored. The spatial proximity of these key residues during the drug-bound state, their accessibility, state dependent or not, from the extracellular and cytoplasmic sides also require additional experiments (e.g., cysteine scanning mutagenesis). For nonfunctional single Ala-substituted channels, the generation of tandem constructs with one, two, or three copies of a mutation introduced the pore may shed insights into their roles in drug block.

To date, four isoforms, namely HCN1–4, have been identified. These isoforms exhibit different patterns of gene expression and tissue distribution [23, 25–28] and coassemble to form heteromeric complexes (except between HCN2 and HCN3) that underlie the native If [11, 36, 38]. Although the primary sequences of the HCN1-4 pores are highly conserved, three of the residues substituted in this study, V356, S357, and V379, are isoform variants (Fig. 1). While S357A only modestly contributes to ZD7288 binding and V356A abolishes channel function (which prevents any direct conclusion about its pharmacological role to be drawn), we have identified V379 as a determinant of drug sensitivity and proposed that it constitutes part of the drug receptor. The isoforms HCN1 and HCN2 are known to display different drug sensitivities. Although residue 379 is an isoform variant, V379I has gating properties and ZD7288 sensitivity not different from WT murine HCN1. Thus, further experiments will be needed to investigate the basis of isoform-specific drug block. In stark contrast, the homologous substitution V390I in human HCN1 markedly increases the ZD7288 sensitivity rendering it HCN2-like [12]. These results suggest that species- as well as isoform-specific differences in drug block exist. Further experiments will be needed to investigate the basis of these differences. Given the fact that native If (or Ih) currents from various tissues have diverse molecular identities due to the different isoforms expressed and their coassembly, a better understanding of drug block of HCN1–4 channels is crucial for developing tissue-selective If blockers.

Fig. 7 Interaction energies of the pairs M377–F378, F378–V379, and M377–V379. a A representative cycle analysis of the interaction pair M377–F378. b Bar graph showing the absolute interaction energies (ΔΔG) for the pairs M377–F378, M377–V379, and F378–V379.
Fig. 8 Schematic diagram showing the proposed model of ZD7288 binding to its receptor constituted by C347, F378, and V379. See text for details.
HCN-encoded $I_f$ (or $I_h$) is one of the key players that prominently modulate the rhythmic firing activity of cardiac nodal pacemaker. For the heart, idiopathic and familial sinus dysfunctions associated with human HCN mutations have been reported [24, 29]. Although cardiac $I_f$ is most abundant in the sino-atrial node, it is also found at various levels in the atrioventricular node, the Purkinje fibers, the atria, and the ventricles [8, 9, 14]. Indeed, upregulation of $I_f$ has been suggested to contribute to atrial ectopy and other arrhythmias associated with heart failure, hypertrophy, and hypertension [8, 9, 14]. As a result, drugs that target HCN channels such as ivabradine, ZD7288, have been developed. Using an engineered HCN1 channel with the S3-ZD7288, have been developed. Using an engineered HCN1 channel with the S3–S4 linker residues 277–279 deleted [33, 34], we have successfully constructed an in vivo bioartificial cardiac pacemaker that suffices to replace or supplement conventional electronic devices in a large animal sick sinus syndrome model [35, 39]. Thus, our present results may be useful for designing future HCN drugs and engineered HCN channels with a custom-tailored modulatory drug receptor for gene- [35, 39] and cell-based [21, 32] therapies. For instance, the substitution M377A can be combined with others to generate a bioartificial pacemaker that is less sensitive to bradycardiac drugs to confer drug specificity and to minimize potential side effects (due to blockade of endogenous $I_f$, or $I_h$ in other tissues).

Of note, the HCN1 isoform examined in the present study is the least sensitive to cAMP among HCN1–4. HCN1 has been chosen because we have previously extensively studied its structure–function properties [1–4, 16, 19, 31, 33, 34, 37, 38]. Indeed, we have even successfully employed an engineered HCN1 channel for constructing bioartificial pacemakers both in vitro and in vivo [10, 22, 35, 39]. Although the least sensitive, HCN1 nonetheless responds to adrenergic stimulations [35]. The present results will also be particularly useful for improving antiarrhythmic designs and selectivity. Nevertheless, further experiments at physiological temperature, though more technically difficult to perform and thus missed in the present study [40], will be useful to examine the temperature-dependent effect as low temperatures was shown to possibly attenuate $I_f$ inhibition [41].

In summary, we conclude that pore residues C347, F378, and V379 of HCN1 channels are crucial for ZD7288 block. Together, these amino acids may constitute part of the drug-binding receptor that is located relatively deep into the pore. These P-loop and the S6 residues exert their effects by modulating drug access to the receptor and by stabilizing the drug-bound HCN channel complex. Therefore, we proposed a refined drug-blocking model that may lead to more improved antiarrhythmics and bioartificial pacemaker designs, in particular, to provide better insights in improving drug selectivity.

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