Direct Measurement of Cardiac Na⁺ Channel Conformations Reveals Molecular Pathologies of Inherited Mutations

Running title: Varga et al.; Cardiac Na⁺ Channel Conformations

Zoltan Varga, PhD¹,²; Wandi Zhu, BS¹; Angela R. Schubert, BS¹; Jennifer L. Pardieck, BS¹; Arie Krumholz, PhD¹; Eric J. Hsu, BS¹; Mark A. Zaydman, BS¹; Jianmin Cui, PhD¹; Jonathan R. Silva, PhD¹

¹Department of Biomedical Engineering, Washington University in St. Louis, St. Louis, MO; ²MTA-DE-NAP B Ion Channel Structure-Function Research Group, RCMM, University of Debrecen, Debrecen, Hungary

Correspondence:
Jonathan R. Silva, PhD
Washington University in St. Louis
Campus Box 1097
1 Brookings Dr.
St. Louis, MO 63130
Tel: 314-935-8837
Fax: 314-935-7448
E-mail: jonsilva@wustl.edu

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Abstract:

**Background** - Dysregulation of voltage-gated cardiac Na⁺ channels (NaV1.5) by inherited mutations, disease-linked remodeling, and drugs causes arrhythmias. The molecular mechanisms whereby the NaV1.5 voltage-sensing domains (VSDs) are perturbed to pathologically or therapeutically modulate Na⁺ current (I_Na) have not been specified. Our aim was to correlate I_Na kinetics with conformational changes within the four (DI-DIV) VSDs to define molecular mechanisms of NaV1.5 modulation.

**Method and Results** - Four NaV1.5 constructs were created to track the voltage-dependent kinetics of conformational changes within each VSD, using voltage-clamp fluorometry (VCF). Each VSD displayed unique kinetics, consistent with distinct roles in determining I_Na. In particular, DIII-VSD deactivation kinetics were modulated by depolarizing pulses with durations in the intermediate time domain that modulates late I_Na. We then used the DII-VSD construct to probe the molecular pathology of two Brugada Syndrome (BrS) mutations (A735V and G752R). A735V shifted DII-VSD voltage-dependence to depolarized potentials, while G752R significantly slowed DII-VSD kinetics. Both mutations slowed I_Na activation, even though DII-VSD activation occurred at higher potentials (A735V) or at later times (G752R) than ionic current activation, indicating that the DII-VSD allosterically regulates the rate of I_Na activation and myocyte excitability.

**Conclusions** - Our results reveal novel mechanisms whereby the NaV1.5 VSDs regulate its activation and inactivation. The ability to distinguish distinct molecular mechanisms of proximal BrS mutations demonstrates the potential of these methods to reveal how inherited mutations, post-translational modifications and anti-arrhythmic drugs alter NaV1.5 at the molecular level.

**Key words:** sodium channels, Brugada syndrome, ion channel
Introduction

Atrial and ventricular action potentials (AP) are initiated by a large membrane-depolarizing Na\(^+\) flux through the cardiac voltage-gated Na\(^+\) channel, Na\(_V\)1.5. In large animals and humans, persistent Na\(_V\)1.5 current is also a significant regulator of AP duration. Because of its central role in determining the AP, molecular-level interactions that perturb Na\(_V\)1.5 channel function have a significant impact on the ability of the myocardium to initiate and sustain arrhythmias.

Mutations to Na\(_V\)1.5 are known to cause Long QT Syndrome Type 3 (LQT3), Brugada Syndrome (BrS), Sick Sinus Syndrome, Atrial Fibrillation and Familial Heart Block \(^{1,2}\). Further, class I small molecule anti-arrhythmics target the Na\(_V\)1.5 pore with the aim of diminishing Na\(^+\) current to prevent arrhythmia\(^3\). To date, characterization of Na\(_V\)1.5 mutation pathology and drug interactions has primarily relied on measuring the ionic current response to elaborately designed voltage-pulse protocols. Here, we develop novel Na\(_V\)1.5 constructs to augment these protocols by monitoring changes in conformation with fluorescence labeling, at the same time that we observe the ionic current, with voltage-clamp fluorometry (VCF). By applying VCF to observe channels that carry inherited mutations, we expect to improve phenotypic precision by characterizing the molecular pathology of the disease.

The Na\(_V\)1.5 \(\alpha\)-subunit, encoded by SCN5A, is composed of four homologous domains (DI-DIV), each with six transmembrane-spanning segments (S1-S6). Within DI-DIV, S1-S4 form the voltage-sensing domains (VSD), which respond to changes in membrane potential \((V_m)\) to cause Na\(_V\)1.5 gating. The Na\(_V\)1.5 pore is formed by the S5-S6 segments. A hydrophobic triplet (IFM) located in the intracellular DIV/DIV linker is required for inactivation\(^4\), and its action is modulated by the C-terminus\(^5\).

Previously, VCF was used to relate skeletal muscle Na\(^+\) channel, Na\(_V\)1.4, VSD...
conformations to channel activation and inactivation\textsuperscript{6-10} and to probe the domain-specific interactions of toxins\textsuperscript{11-13} and local anesthetics\textsuperscript{14, 15}.

In this study, we present four novel constructs that report on the distinctive VSD conformations of Na\textsubscript{V}1.5 for the purpose of assessing molecular mechanisms of Na\textsubscript{V}1.5-linked pathologies and therapies. To demonstrate the potential of the method, we probe the molecular phenotype of two proximal BrS1 mutations, A735V and G752R, which are both located on the DII-VSD. The ECG phenotypes of patients with each mutation differ. Patients with A735V present with S-T segment elevation in leads V1-V3\textsuperscript{16}, while those with G752R show S-T segment elevation\textsuperscript{17} in addition to a prominent J wave in leads II, III, and aVF\textsuperscript{18}. We hypothesized that in addition to ECG differences, there may also exist fundamentally different molecular pathologies, despite the proximity of the mutations.

Materials and Methods

Voltage clamp fluorometry
cRNAs for human Na\textsubscript{V}1.5 \(\alpha\) and \(\beta_1\)-subunits were injected into \textit{Xenopus laevis} oocytes for cut-open oocyte recordings\textsuperscript{19, 20} at 19°C. The internal solution was (mmol/L): 105 NMG-Mes, 10 Na-Mes, 20 HEPES, and 2 EGTA, pH 7.4, and the external solution was composed of (mM): 25 NMG-Mes, 90 Na-Mes, 20 HEPES, and 2 Ca-Mes\textsubscript{2}, pH 7.4. For gating currents, 10 \(\mu\)M TTX was used, and Na-Mes was replaced by NMG-Mes. For fluorescence recordings, oocytes were labeled with 10 \(\mu\)mol/L methanethiosulfonate-carboxytetramethylrhodamine (MTS-TAMRA). Data was collected on a custom rig (see \textit{Supplemental Methods}). Animal procedures were performed according to institutional guidelines.

Data analysis

For photobleaching correction, baseline fluorescence recorded over time without voltage pulsing
was subtracted from traces recorded during the application of the voltage protocol. Fluorescence magnitude is expressed as \( \Delta F/F_0 \), \( \Delta F \) is the signal amplitude change and \( F_0 \) is baseline fluorescence.

Steady-state voltage-dependence was quantified by Boltzmann-function fitting:
\[
y = \frac{1}{1 + \exp\left(\frac{V-V_{1/2}}{k}\right)}
\]
t_{10-90\%} for current activation is the duration between 10 and 90% of the ionic current peak. For fluorescence signals, \( t_{10-90\%C} \) was calculated from the time constants of single exponential fits (\( \tau \)).

Statistical comparisons were done using paired or independent-sample t-tests or one-way ANOVA with post-hoc tests for pairwise comparison. \( p<0.05 \) was considered significant. The ± symbols in the text and error bars in the figures represent the 95% confidence interval (95CI) with number of trials (n) in parentheses.

**Results**

Our first aim was to create four Na\(_V\)1.5 constructs to track conformational changes within each of the Na\(_V\)1.5 channel VSDs. Initial attempts to record fluorescence signals were unsuccessful due to insufficient expression of human Na\(_V\)1.5, which is significantly reduced compared to rat Na\(_V\)1.4, the isoform used to acquire most existing Na\(^+\) channel VCF data. Examination of differences between the two channels revealed that Na\(_V\)1.5 contains an ubiquitination motif that enhances the channel recycling rate and reduces the number of channels in the membrane\(^{21}\); this motif is not found in Na\(_V\)1.4. Ablation of the ubiquitination site by the Y1977A mutation and co-injection with \( \beta_1 \)-subunit RNA resulted in sufficient Na\(_V\)1.5 expression for the detection of useful fluorescence signals. We also introduced C373Y, increasing tetrodotoxin\(^{22}\) (TTX) sensitivity and allowing ionic current blockade for the measurement of gating currents. This mutation also removes an externally accessible cysteine, preventing non-specific labeling.
Two protocols were performed to compare our new construct to wild type. We first quantified the conductance-voltage (G-V) relationship, found by measuring peak current after a series of depolarizing pulses and dividing by driving force. The second protocol measured the voltage-dependence of steady state inactivation (SSI); peak current was measured during a short test pulse to -20 mV that followed 200-ms inactivation-inducing pulses to varying potentials. The voltage-dependence of the G-V and SSI curves were not substantially affected by these mutations (Fig. 1, p=0.951 and 0.069, respectively). For fluorescence tracking, cysteine mutations (specified below) in the individual domains were introduced in this Y1977A-C373Y mutant, which we named WT-LFS (Large Fluorescence Signal). The mutants produced for VSD-tracking in this background will be accordingly referred to as DI-LFS, DII-LFS, DIII-LFS, and DIV-LFS. Cysteines were introduced at several positions within each VSD, but most of these did not produce useful signals. We were fortunate to find a single useful site for each domain.

To track the conformation of the DI-VSD, we tested ten positions from L212 to T220 and R222. V215C (DI-LFS) was the only mutant to express well and produce a usable fluorescence signal (Fig 2). The Na\(_V\)1.5 VSDs play an essential role in determining gating, and we expected the TAMRA-MTS labeling in conjunction with the introduced cysteine to affect gating to some extent. We observed a negative shift (\(\Delta V_{1/2}=-16.8\) mV) in the G-V and a slight positive shift (\(\Delta V_{1/2}=+8\) mV) in the SSI curves compared to the background WT-LFS construct, implying that the labeled V215C channel opens at more negative potentials and inactivates at more positive potentials than WT-LFS. Comparing labeled and unlabeled DI-LFS showed that the G-V shift is a consequence of adding the TAMRA-MTS label (Figure S1). We also used gating currents to assess whether charge movement across the transmembrane electric field was affected by labeling (Fig. S2). If the probe significantly affected charge transfer, we would expect to
observe a change in the voltage-dependence of the gating current integral, the gating charge. For the four constructs, the charge-voltage (Q-V) relationship was shifted over a range of 5 to 11 mV in comparison to WT-LFS (Fig. S1 and Table 1), suggesting that charge displacement was modestly affected by cysteine-conjugated fluorophore replacement of the WT residue.

Simultaneously recording changes in fluorescence magnitude during these protocols allowed us to correlate DI-VSD conformations to channel gating. We refer to the fluorescence that was recorded during the G-V protocol as the fluorescence-voltage relationship (F-V curve) and the fluorescence from the SSI protocols as the SSI F-V curve. The F-V curves reflect the voltage-dependence of the conformations in the DI-VSD that are reported by TAMRA. For the DI-VSD, the midpoint of the F-V curve was 27.2 mV negative of the G-V curve, indicating that DI-VSD movement occurs at much more negative potentials than pore opening. On the contrary, the SSI and SSI F-V curves had very similar midpoints, showing that inactivation and DI-VSD movement occur in the same voltage range. However, in contrast to DIV (see below), the DI-VSD SSI F-V and SSI slopes differed substantially (k=-15.4(5) vs -6.4 mV(8), respectively, p<0.001), suggesting that the two events are not tightly coupled.

The fluorescence kinetics recorded during the F-V curve reflect the voltage-dependent rates of VSD activation and deactivation. Since current and VSD activation kinetics are described by different functions, we measured the time between 10% and 90% of the peak current and calculated t_{10-90\%} for fluorescence (see Methods). DI-VSD rise-time was slower (t_{10-90\%}=0.82±0.23 ms at +30 mV) than ionic current activation (t_{10-90\%=}0.24±0.04 ms), but the events overlapped due to the sigmoidal delay in ionic current activation (Fig. S3). At elevated potentials (> +20 mV) a slow component in the DI-VSD fluorescence signal became apparent with an opposite sign relative to the original deflection. While significant numbers of channels
will not make this slow transition to a new state during a single AP, channels may accumulate in this state at rapid heart rates.

Since fast inactivation is known to hinder the recovery of the gating charges upon repolarization (immobilization), we performed experiments to measure the deactivation rate of the DI-VSD after depolarizing pulses of varying length, which inactivated the channels to different extents. The deactivation kinetics measured at -120 mV did not change as a function of pulse duration (t$_{10-90\%C}$=0.99±0.22 ms after a 2 ms pulse and 0.99±0.13 ms after a 200 ms pulse, (4), p=0.948). This result suggests that DI-VSD deactivation is not influenced by NaV1.5 open-state inactivation, which had not occurred during the 2 ms pulse and is nearly complete after 200 ms.

We tracked the DII-VSD with S805C (DII-LFS), which yielded large and robust signals. The voltage-dependent steady-state parameters (G-V and SSI curves) were minimally affected in comparison to WT-LFS; there was no shift in the G-V curve and a +7.4 mV shift in the SSI curve (Fig. 3). Measurement of the F-V revealed that it was 12.5 mV negative of the G-V curve, showing that the DII-VSD starts to move prior to activation; however, it reaches saturation at higher potentials than the conductance due to its shallower slope, which indicates that the two events are not tightly coupled. Comparing the SSI and SSI-FV curves shows that inactivation occurs at more negative potentials than DII-VSD activation (V$_{1/2}$=-79.2 and -52.2 mV, respectively), and that the inactivation slope is much steeper than DII-VSD activation (k=-6.5 and -17.5 mV, respectively, (7,6), p<0.001). Thus, at holding potentials which completely abolish the conductance by inactivation (V>-50 mV), the DII-VSD is still far from having completed its motion, likely precluding it from participating in closed-state inactivation. The DI-VSD and DII-VSDs activate at similar rates (t$_{10-90\%C}$=0.98±0.19 ms at +30 mV, (5) p=0.332),
and, when superimposed on the ionic current, DII-VSD activation precedes ionic current activation (Fig. S3). As for DI, the DII-VSD deactivation kinetics are not substantially affected by the length of the depolarizing pulse ($t_{10-90\%C} = 0.86 \pm 0.29$ ms after 2 ms and $0.99 \pm 0.13$ ms after 200 ms, (4), p=0.171).

Conformations in the DIII-VSD were tracked with M1296C (DIII-LFS), which was the only construct to produce a signal out of I1299C, K1300C, and S1301C. The DIII construct had G-V and SSI curves nearly identical to those of DII, but relative to the DII-VSD, the F-V and SSI F-V curves were dramatically shifted by -54.6 and -70.3 mV, respectively (Fig. 4). Comparison to the G-V and SSI curves shows that DIII-VSD activation precedes pore opening by a large voltage margin during activation, and similarly, during inactivation, most of the VSD movement is completed before any noticeable drop in the conductance due to inactivation.

Activation kinetics of the DIII-VSD were similar to those of DI and DII ($t_{10-90\%C} = 0.81 \pm 0.25$ ms at +30 mV, (5), p=0.332), although deactivation was >10x slower. Furthermore, in contrast to DI and DII, the deactivation kinetics of the DIII-VSD were significantly slowed when the duration of depolarizing pulse was increased ($t_{10-90\%C} = 12.85 \pm 2.37$ ms after a 2 ms pulse and $18.58 \pm 1.58$ after 200 ms, (6), p=0.009). The markedly slowed deactivation of the DIII-VSD following long depolarizations indicates that its return could be inhibited by entry of the channel into the deeper inactivated states that occur after longer depolarizations, which stabilize it in the activated conformation.

Finally, to track the DIV-VSD, we introduced S1618C (DIV-LFS) and L1621C (no signal). The unlabeled DIV-LFS construct displays a non-saturating SSI curve that is corrected by fluorophore conjugation. Recording of the unlabeled construct with a reducing agent confirmed that this phenomenon was due to disulphide bond formation in the absence of the
fluorophore (Figure S1). Both the G-V and SSI curves of the DIV-VSD construct were shifted relative to WT-LFS by +5.3 and +12.4 mV, respectively (Fig. 5). Comparison of the fluorescence and ionic current shows that the midpoint of the F-V curve was 32.2 mV more negative than that of the G-V, while the SSI and SSI F-V curves overlapped, further supporting strong correlation between DIV-VSD activation and closed-state inactivation that is primarily measured by the SSI curve. In addition, slowing of fast inactivation caused by the introduction of the cysteine mutation is also readily apparent in the ionic current recordings. In comparison to the other three domains, the DIV fluorescence signal activated much more slowly ($t_{10\%C}=1.91\pm0.44$ ms at $+30$ mV, (7), $p<0.001$). Deactivation kinetics were significantly slower than those of the DI and DII, but 2-3 times faster than that of DIII. Surprisingly, DIV-VSD deactivation did not slow with increasing pulse duration ($t_{10\%C}=5.71\pm4.04$ ms after a 2 ms pulse and $6.63\pm3.91$ ms after 200 ms, (4), $p=0.411$). Given the prominent role of DIV in inactivation, the lack of slowing in DIV-VSD recovery with longer pulses is highly unexpected and is in clear contrast with NaV1.4 results$^8$, suggesting different interactions between the DIII- and DIV-VSDs and inactivation.

Our next aim was to assess the molecular pathology of two proximal DII-VSD BrS mutations, A735V and G752R (Fig. 6). We introduced each into the DII-LFS construct. The F-V and SSI F-V curves of A735V were dramatically positive-shifted relative to DII-LFS (by +49.1 mV and +58.5 mV, respectively) and also relative to the G-V and SSI curves of the channel itself (by +19.5 mV and +89.1 mV, respectively). In addition, the voltage-sensitivity of the DII-VSD was severely reduced; both had very shallow slope factors (35.4 mV and -39.5 mV, respectively). DII-VSD activation kinetics were also slightly slowed by the mutation at +40 mV based on the $t_{10-90\%C}$ values (DII-LFS: $0.99\pm0.19$ ms (8), DII-LFS-A735V: $1.49\pm0.36$ ms, (5),
p=0.005), but even more strikingly, the DII-VSD activation rate of the mutant lost its voltage-dependence (1.45±0.42 ms at -20 mV and 1.49±0.36 ms at +40 mV (5), p=0.850) as opposed to DII-LFS (1.42±0.19 ms at -20 mV and 0.99±0.19 ms at +40 mV (8), p=0.002). To the contrary, ionic current activation of the mutant conserved its voltage-dependence, but was slower than DII-LFS at potentials <10 mV (t_{10-90%}=1.48±0.17 ms (5) vs 0.71±0.47 ms (3), respectively, at -20 mV, p=0.002). Despite the severely right-shifted F-V of A735V, the mutant Q-V (V_{1/2} = -75.6±8.3 mV (5)) was not significantly different from that of DII-LFS (-80.0±7.6 mV (4), p=0.241) or WT-LFS (-69.1±4.2 mV (7), p=0.058).

In contrast, G752R caused minor F-V and SSI F-V shifts (by +1.0 mV and -9.8 mV, respectively, compared to DII-LFS) and displayed similar slope factors, but resulted in a severe slowing of the DII-VSD activation. In response to a depolarizing step, the rate of DII-VSD activation was about 1-fold slower in the mutant than in the DII-LFS channel (t_{10-90%}=0.99±0.19 ms (8) and 6.96±1.21ms (8) at +40 mV, respectively, p=0.001). The Q-V curve was right-shifted by ~20 mV with respect to DII-LFS. However, extending the interval over which charge was integrated from 10 to 20 ms, to account for G752R-induced slowing, shifted the curve leftward and closer to DII-LFS. Significant slowing of activation kinetics was also observed in the ionic current of G752R. In the -40 to +30 mV voltage range, the activation rate was 1.7 to 2.9 times slower than that of the DII-LFS as measured by the t_{10-90%} times (1.51±0.15 ms (7) and 0.71±0.47 ms (3), respectively, at -20 mV, p<0.001; Fig S4).

To ensure that the mechanisms we observed in oocytes are still operative across cell lines, we expressed both mutants in HEK293T cells. A comparison (Fig. S5) shows that as in oocytes, current activation measured by the t_{10-90%} was slowed in both BrS mutants compared to DII-LFS (at -20 mV, DII-LFS: 0.27±0.09 ms (4), A735V: 0.81±0.32 ms (5), G752R: 0.66±0.07
Discussion

Our results represent the first measurements of human NaV1.5 VSD conformations and show that each VSD has unique kinetics, consistent with specific gating roles. Specifically, the four VSDs activate at very different membrane potentials that span a range >50 mV. DIII activates at the most hyperpolarized potentials (V_{1/2}=-106.0 mV), followed by DI(-83.9 mV), DIV(-66.8), and finally DII(-51.4), all negative of the G-V curve that characterizes current activation (V_{1/2}=-39.9 mV for WT-LFS). Further, all four VSDs have a t_{10-90} rise time that is slower than ionic current activation, which is to be expected if ionic current activation is due to the combined motions of several VSDs that can be modeled as a cubed gate (e.g., m^3 in Hodgkin-Huxley type models), allowing for a more rapid activation after a delay.

While our results support the hypothesis of concerted action of several VSDs in activating NaV channels, our mutation results contrast with classical models that require tight coupling of the “gates” to pore opening. Comparing DII-VSD activation voltage-dependence in A735V to its G-V relationship (Fig. 6A) reveals that the channel fully opens at 0 mV, even though the DII-VSD is not fully activated even at +80 mV. This phenomenon is observed in DII-LFS as well, indicating that this property is intrinsic to NaV1.5, but the effect is greatly enhanced in A735V. Thus, DII-VSD activation facilitates channel opening through an allosteric mechanism where lack of VSD-activation simply slows activation, but does not prevent it. This mechanism may be evolutionarily beneficial, because it allows for moderate alteration of VSD kinetics by mutations without preventing Na^+ channel activation and myocyte excitability.

Correlation of VSD-activation with inactivation can be assessed by comparing the SSI and SSI-FV curves and pulse duration protocols. The SSI protocols show that the DII-VSD is
least likely to participate in inactivation: At potentials when the channel is completely blocked by inactivation, the DII-VSD has yet to significantly activate (Fig. 3). In contrast, the close overlap of the SSI and DIV SSI F-V curves supports the notion that the closed-state inactivation reported by this protocol is tightly coupled to DIV-VSD movement. Finally, the pulse duration protocol may reveal VSD interaction with deeper inactivation states that occur at depolarized potentials, which are likely to modulate the late Na\(^+\) current.

**The origin of \(\Delta F\)**

Our detected changes in TAMRA fluorescence emission are likely due to photo-induced electron transfer, which is a quenching non-radiative electron transfer between two molecules that are within 10 Å, in close van der Waals contact\(^{23}\). Many different residues may quench the fluorophore, which implies that our fluorescence signal is highly dependent on the surrounding residues that interact with the fluorophore as it moves. This dependence introduces the possibility that the fluorescence changes we observe do not necessarily reflect the movement of the charged S4 segments across the membrane.

Our Q-V curves that are calculated from gating currents directly quantify charge transfer within all four domains across the membrane field. Thus, we expect that if we are tracking charge transfer, then the valence-weighted average midpoint of the F-V curves will match the Q-V curves (Figure 7, Table 1)\(^6\). The WT-LFS Q-V is accurately reconstructed by the valence-weighted F-V curves (Figure 7A). Moreover the confounding left-shift in the A735V Q-V relative to G752R is also predicted by this analysis (Figure 7B), and results from much less charge being assigned to the DII F-V due to its reduced slope, which lessens its right-shifting contribution. With the G752R construct, we observe less charge transferred at some of the lower potentials. This slight difference is likely due to the substantial slowing of charge...
movement at these potentials, which sharply reduces the gating current and precludes its detection. Alternatively, G752R may be slowing activation of the other three domains, causing a right-shifted Q-V. Previous work in Na\textsubscript{V}1.4 shows that a DII-VSD activation shift modestly affects the activation of the DI, DIII, and DIV VSDs\textsuperscript{25}. Future efforts to probe Na\textsubscript{V}1.5 cooperativity will be needed to test whether this the case for A735V and G752R.

A simple two-state model of the VSD would predict that the SSI-FV is simply a mirror image of the F-V and that they should cross at 0.5. This is the case for the DI and DII VSDs, but not for the DIII- and DIV-VSDs, which showed a lower crossing. Since the SSI-FV uses a 200 ms pulse, whereas the F-V is measured after several ms, the difference is likely due to multiple activated or resting states of the VSD. This is not surprising, as multiple DIII-VSD resting states have been posited to model lidocaine interaction and we also infer multiple states from the pulse duration protocol. For the DIV-VSD this crossing was in the range of experimental variability, but may be linked to DIV-VSD interaction with inactivation.

**Comparison to Previous Results**

VCF has been used to study Na\textsubscript{V}1.4, and many of the results have been extrapolated across the super-family of Na\textsubscript{V} channels. While we observe many similarities, there are several distinct Na\textsubscript{V}1.5 features. In rNa\textsubscript{V}1.4, the VSD activation midpoints all fell within a 20 mV range. In contrast, midpoints of the F-V curves for hNa\textsubscript{V}1.5 spanned a range of about 50 mV. In particular, DIII was activated at very hyperpolarized potentials, and DII activation was substantially depolarized relative to the Q-V (by 30 mV). For the DII-VSD, this depolarizing shift may be due to loose coupling between the DII-VSD and channel activation, which was not observed in Na\textsubscript{V}1.4.

Further differences are seen when comparing the connection between inactivation and
VSD activation. Classical studies of Na\textsubscript{V} channel gating charge movement found that, following prolonged depolarization, a fraction of the gating charge is “immobilized”: that is, it returns with much slower kinetics upon repolarization than the majority of the charge\textsuperscript{26}. In human Na\textsubscript{V}1.4, this immobilized fraction was associated with DIII and DIV\textsuperscript{8}. Consistently, hNa\textsubscript{V}1.5 DI- and DII-VSD deactivation was independent of the length of the depolarizing pulse. Conversely, return of the hNa\textsubscript{V}1.5 DIII-VSD was significantly slowed by long depolarizations, while the DIV-VSD deactivation rate was unaffected, in clear contrast to Na\textsubscript{V}1.4. However, this observation is in agreement with previous reports showing that charge immobilization of the Na\textsubscript{V}1.5 DIV-VSD is independent of inactivation, and that the inactivation particle can only modulate gating charge recovery of DIII, but not DIV\textsuperscript{27}. Thus our data reconciles these two previously discrepant results by showing that the hNa\textsubscript{V}1.5 VSDs interact with inactivation uniquely.

Many have previously probed the functional contributions of each VSD to Na\textsubscript{V}1.5 gating, and our data shows the timing of these contributions. For example, outward stabilization of the DI-S4 strongly reduced peak Na\textsubscript{V}1.5 current (45%) without affecting rapid channel kinetics\textsuperscript{28}. In our data, we observed a slow DI-VSD transition at elevated potentials that may be linked to this peak reduction through slow inactivation (Figure 2). In contrast, DII-VSD outward stabilization does affect rapid gating kinetics, shifting the G-V leftward along with SSI\textsuperscript{28}. Our results show that this DII-VSD regulation of channel activation is through an allosteric connection, which facilitates channel opening when the DII-VSD is activated. Similar work has also probed the consequences of DIII and DIV-VSD outward stabilization\textsuperscript{29}, both of which alter channel availability without affecting activation. While the link between the four VSDs and inactivation is consistent with our results, it remains to be seen whether a slowing of DI, DIII or DIV may
also affect channel activation as observed in DII.

Relevance

For the inherited BrS1 mutations, A735V and G752R, we observed two very different molecular pathologies (Fig 6). A735V clearly shifts the voltage dependence, while at positive potentials DII-VSD activation kinetics are nearly as rapid as WT. In contrast, DII-VSD voltage-dependence is preserved with G752R, but it arrives at its steady state much more slowly. For both mutants, the consequence for the ionic current is slower activation. Thus, our results show that dramatically different molecular pathologies can give rise to very similar ionic current phenotypes. Still, molecular-level differences are likely to have significant consequences for the interaction of the channel with anti-arrhythmic therapies.

From a theoretical perspective, the non-canonical Na_V1.5 activation mechanism revealed by the G752R and A735V mutations suggests a very specific Markov state diagram where opening is possible from closed states where the VSDs are not yet activated. Recent work on L-type Ca^{2+} channels revealed that this type of allosteric model was best suited to describe its data. Given the recent attention to similarities between Ca^{2+} and Na^{+} channels it would not be surprising if a similar model would be more suitable for describing Na_V1.5 gating. We expect that future work probing the contribution of each VSD to channel gating will be immensely useful for informing these widely used models.

Future Directions

As demonstrated by our ability to observe the molecular pathology of two BrS1 mutations, we expect that we will be able to better understand mechanisms whereby some of the >150 Na_V1.5 inherited mutations cause disease. In addition, Na_V1.5 is known to interact with many accessory proteins, and it is tightly controlled by post-translational modification. By observing how these
modulatory proteins and modifications affect the VSDs, we expect to unravel molecular mechanisms whereby the cell regulates hNav1.5 gating to modulate cardiac myocyte excitability.

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**Conflict of Interest Disclosures:** None.

**References**


7. Silva JR, Goldstein SA. Voltage-sensor movements describe slow inactivation of voltage-


Table 1: Parameters of Boltzmann-fits to G-V, F-V, SSI, SSI-FV and Q-V curves

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Errors represent 95% confidence intervals, numbers in parentheses indicate the number of cells measured.
Figure Legends:

Figure 1: WT-LFS ionic and gating currents. Currents from WT-LFS and WT channels were recorded in the cut-open oocyte configuration. The mean±95CI for groups of 6 to 22 cells are reported. (A) Top: Ionic currents. From -100 mV, a 50 ms step to -120 preceded 100 ms depolarizing pulses from -140 to +40 mV in 10 mV steps, which were followed by a 50 ms step to -120 mV before returning to holding potential. Membrane capacitance and leak were removed using P/-8 leak subtraction. For clarity, only the first 40 ms of traces corresponding to -130 to +30 mV in 20 mV steps are shown. Bottom: Voltage-dependence of steady-state activation and steady-state inactivation curves for the WT-LFS (G-V: black circles, SSI: black squares) and the WT channels (G-V, gray up triangle; SSI, gray down triangle). G-V curves were constructed from I-V relationships recorded from cells depolarized in 10 or 20 mV increments from -120 mV. Na\(^+\) reversal potential was determined for each cell individually. For the SSI curve, cells were held at potentials ranging from -150 to +20 mV for 200 ms, and the available fraction was assessed by a -20 mV test pulse. See Table 1 for Boltzmann-fit parameters. (B) Top: Gating currents from WT-LFS channels were recorded during 20-ms depolarizing steps ranging from -150 to +50 mV in 20-mV steps. Capacitance and leak were removed using P/4 leak subtraction with a sub-sweep holding level of +40 mV. Bottom: Gating currents were integrated for 10 ms following the voltage change.

Figure 2: V215C (DI-LFS) ionic currents and fluorescence. Na\(^+\) currents and fluorescence were recorded from human Na\(_V\)1.5 channels carrying the LFS-S215C mutations with cut-open oocyte voltage-clamp fluorometry, see Materials and Methods. The mean±95CI for groups of 4 to 14
cells is reported. A) Top: Ionic currents (above) and fluorescence signals (below). From -100 mV, a 50 ms step to -120 preceded 20 ms-long depolarizing pulses ranging from -150 to +30 mV in 20 mV steps. These steps were then followed by a 50 ms step to -120 mV before returning to the holding potential. Membrane capacitance and leak were removed using P/-8 leak subtraction. For clarity, only traces corresponding to -150, -90, -30, and +30 mV are shown. Bottom: G-V curve (black circles) and SSI (black squares) and the corresponding fluorescence signals (orange circles and squares, respectively). All curves were obtained by Fig. 1 protocols. The fluorescence amplitude for the FV and SSI-FV curves was determined by taking the mean of the signal for at least 3 ms at the greatest displacement of the signal. For the SSI-FV curve, ΔF was calculated as the difference between the signal amplitude measured at the conditioning potential and -20 mV. Boltzmann-fit parameters in Table 1. Dotted lines represent WT-LFS G-V and SSI curves for comparison. B) Top: Tail currents and fluorescence measured at -120 mV following depolarizing steps to +50 mV for 2 (black) or 200 ms (orange). Inactivation does not occur during the shorter pulse resulting in a large tail current. Bottom: t10,90%,C values calculated from deactivation time constants of the fluorescence signal from exponential fits following the step back to -120 mV, from depolarizing steps to 0 or +50 mV for 1 to 200 ms durations.

**Figure 3:** DII-LFS (S805C) ionic currents and fluorescence. Na+ currents and fluorescence were recorded from DII-LFS. Conditions were as in Figure 2 unless noted. Mean±95CI is reported for groups of 3 to 9 cells. A) Top: Ionic currents (above) and fluorescence (below). Bottom: G-V curve (black circles), SSI (black squares), and the corresponding fluorescence (green circles and squares, respectively). Curves were constructed as in Figure 2. Boltzmann-fit parameters in Table 1. Dotted lines represent WT G-V and SSI curves for comparison. B) Tail currents and
fluorescence measured following the step back to -120 mV from depolarizing steps to +50 mV for 2 (black) or 200 ms (green). $t_{10-90\%}$ values calculated from deactivation time constants of the fluorescence signal from exponential fits following the step back to -120 mV, from depolarizing steps to 0 or +50 mV for 1 to 200 ms durations.

**Figure 4:** DIII-LFS (M1296C) ionic currents and fluorescence. Na$^+$ currents and fluorescence from DIII-LFS. Mean±95CI is reported for groups of 5 to 17 cells. A) Top: Ionic currents (above) and fluorescence (below) were recorded during 20 ms pulses ranging from -170 to +30 mV in 20 mV steps. For clarity, only traces corresponding to -170, -110, -50, and +10 mV are shown. Bottom: G-V (black circles), SSI (black squares), and the corresponding fluorescence signals (blue circles and squares, respectively). Curves were constructed as in Figure 2. Boltzmann-fit parameters in Table 1. For comparison, dotted lines represent WT G-V and SSI curves. B) Tail currents and fluorescence measured following the step back to -120 mV from depolarizing steps to +50 mV for 2 (black) or 200 ms (green) pulses. Red dotted lines indicate exponential fits to the fluorescence traces. Time constants for this cell were 4.1 and 7.5 ms after 2 and 200 ms-long depolarizations, respectively. $t_{10-90\%}$ values calculated from deactivation time constants of the fluorescence signal from exponential fits following the step back to -120 mV, from depolarizing steps to 0 or +50 mV for 1 to 200 ms durations.

**Figure 5:** DIV-LFS (S1618C) ionic currents and fluorescence. Na$^+$ currents and fluorescence from DIV-LFS. Mean±95CI is reported for groups of 3 to 20 cells. A) Top: Ionic currents (above) and fluorescence (below) as in Figure 2. Only traces from -150, -90, -30, and +30 mV are shown for clarity. G-V (black circles) and SSI (black squares), and the corresponding
fluorescence signals (purple circles and squares, respectively). Curves were constructed as in Figure 2. Boltzmann-fit parameters in Table 1. Dotted lines represent WT G-V and SSI curves for comparison. B) Top: Tail currents and fluorescence measured during step back to -120 mV from steps to +50 mV for 2 (black) or 200 ms (green) pulses. Bottom: t10-90%,C values calculated from deactivation time constants of the fluorescence signal from exponential fits following the step back to -120 mV, from depolarizing steps to 0 or +50 mV for 1 to 200 ms durations.

**Figure 6:** DII-LFS-A735V and DII-LFS-G752R Brugada syndrome mutants. Na⁺ currents and fluorescence were recorded from A735V (left panels) and G752R (right panels) mutations in the DII-LFS background. Mean±95CI is reported for groups of 3 to 8 cells. Ionic currents (top) and fluorescence (below) from DII-LFS-A735V (A) and DII-LFS-G752R (B) channels were recorded during 50 ms-long pulses ranging from -140 to +60 mV in 20 mV steps. For clarity, only -140, -80, -20 and +40 mV traces are shown. Kinetics of VSD activation in DII-LFS-A735V and DII-LFS G752R channels are tracked by the fluorescence signal. For comparison the gray dotted line represents a normalized signal from DII-LFS recorded at +40 mV. G-V (black circles) and SSI (black squares), and the corresponding fluorescence signals (green circles and squares, respectively) for DII-LFS-A735V and DII-LFS-G752R channels. For comparison, dotted lines represent DII-LFS curves. Voltage-dependence of integrated gating charge movement over 10 ms for A735V and G752R channels. Dashed black line indicates the Q-V function of the ON gating current of WT-LFS for reference. For G752R, dashed green line shows gating charge calculated from a 20 ms interval.

**Figure 7:** Comparison of valence-weighted F-V functions to Q-V functions. F-V curves from
each domain and the valence-weighted sum of these functions\textsuperscript{24} (grey) are shown along with the Q-V curves (black) for (A) WT-LFS, (B) A735V and (C) G752R. For G752R, the 20 ms integral is shown (Fig 6B), and the F-V was also measured at 20 ms.
Direct Measurement of Cardiac Na\textsuperscript{+} Channel Conformations Reveals Molecular Pathologies of Inherited Mutations
Zoltan Varga, Wandi Zhu, Angela R. Schubert, Jennifer L. Pardieck, Arie Krumholz, Eric J. Hsu, Mark A. Zaydman, Jianmin Cui and Jonathan R. Silva

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SUPPLEMENTAL MATERIAL

Supplemental Methods

Molecular Biology
All mutagenesis was performed using the overlap extension polymerase chain reaction (PCR) with Phusion Hot Start Flex Kit (New England Biolabs) and primers from Sigma-Aldrich. PCR products were ligated into the parent plasmid using T4 DNA ligase (New England Biolabs) and were chemically transformed into Turbo Competent cells (New England Biolabs). Multiple clones were selected and plasmids were isolated using the NucleoSpin Plasmid miniprep kit (Macherey-Nagel). Samples were then sequenced to select a single clone for Midiprep (NucleoBond Xtra Midi, Macherey-Nagel). Each construct was then linearized and purified with the NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel). Finally, capped mRNA was synthesized (mMessage mMACHINE T7 Transcription Kit, Life Technologies).

Xenopus laevis
Oocyte harvest was performed on Xenopus laevis (Nasco) no more than four times (~25 frogs). During the oocyte harvest procedure, distress was minimized by anesthetizing frogs in water containing 0.2% tricaine (3-aminobenzoic acid ethyl ester, methane sulfonate salt). After oocyte harvest, the wound was sutured while the frog is still anesthetized, according to our protocols that were approved by the Institutional Animal Care and Use Committee and in accordance with NIH guidelines.

Cut-open oocyte recording
cRNAs for the human β3 subunit (UniProtKB/Swiss-Prot accession no. Q07699.1) and α-subunit NaV1.5 (accession no. Q14524.1) were produced from the pBSTA and pMAX vectors (respectively) and injected at a 2:1 molar ratio (50 ng per cell total) into Xenopus laevis oocytes. Oocytes were incubated at 18°C for 2–7 d in solution with (mM) 93 NaCl, 5 KCl, 1.8 CaCl₂, 1 MgCl₂, 5 HEPES, and 2.5 Na-pyruvate, and 1% penicillin-streptomycin, pH 7.4. Cut-open oocyte recordings were performed using an amplifier (CA-1B; Dagan Corporation) coupled to an A/D converter (Digidata 1440; Molecular Devices) with Clampex and Clampfit software (v10; Molecular Devices) for acquisition and analysis. Temperature was maintained at 19°C with a controller (HCC-100A; Dagan Corporation). The internal solution was (mM): 105 NMG-Mes, 10 Na-Mes, 20 HEPES, and 2 EGTA, pH 7.4. The external solution was composed of (mM): 25 NMG-Mes, 90 Na-Mes, 20 HEPES, and 2 Ca-Mes₂, pH 7.4. For gating currents, Na-Mes was replaced by NMG-Mes in both the external and internal solutions and the external solution contained 10 µM TTX.

Data was collected with standard I-V protocols by simultaneous recording of the ionic current and the fluorescence signal from the labeled domain to construct the voltage dependence of steady-state activation (G-V) and voltage dependence of VS movement (F-V) functions. Cells were depolarized in 10 or 20 mV increments from a holding potential of -100 mV. Depolarizing pulses were preceded by a 100-ms-long prepulse and 50-ms-long postpulse to -120 mV. Capacitance compensation and P-/8 leak subtraction were applied prior to recording ionic currents. To obtain the voltage-dependence of steady-state inactivation (SSI) and the corresponding VS movement (SSI F-V) curves cells were held at the conditioning potential for 200 ms and the available channel fraction was assessed by a depolarizing pulse to -20 mV. For gating currents P/4 subtraction was used with an interpulse holding potential of +40 mV.

VCF measurements
Oocytes were labeled with 10 µM methanethiosulfonate-carboxytetramethylrhodamine (MTS-TAMRA; Santa Cruz Biotechnology) in a depolarizing solution (mM: 110 KCl, 1.5 MgCl₂, 0.8 CaCl₂, 0.2 EDTA and 10 HEPES, pH 7.1) on ice for 20 min. MTS-TAMRA stock solution was 10 mM in DMSO and stored at -20°C. Illumination was provided by a green, high-powered LED (Luminus, PT-121) controlled through a driver
For attached data references to function: photodiode The measurements for points t10 ... noise amplification of the photocurrent. The fluorescence emission was focused onto the photodiode active area using an achromatic doublet (Thorlabs Inc.) with a focal distance of 25 mm.

Data analysis
For the analysis of fluorescence data, baseline fluorescence traces were recorded with no change of voltage during the illumination period. To correct for photobleaching, this baseline trace was filtered and subtracted from the fluorescence traces recorded during the application of the voltage protocol. The magnitude of fluorescence signals is expressed as ΔF / F0, where ΔF is the change in the signal amplitude in response to the voltage change and F0 is the baseline fluorescence.

Steady-state G-V, SSI, F-V and SSI F-V curves were obtained by fitting the data points with a Boltzmann-function: y = 1 / (1 + exp[-(V-V1/2)/k]), where V1/2 is the half-activation voltage and k is the slope factor. For direct comparison of current and VSD kinetics we used t10-90% due to the different functions they follow. t10-90% for current activation kinetics was determined by the time duration between the time points when the current reached 10 and 90 % of the peak value. For fluorescence signals with low signal to noise ratio t10-90% determination was inaccurate, therefore for all fluorescence signals we calculated t10-90% from the time constants of single exponential fits (τ) to the signal by t10-90% = 2.197τ.

Valence-weighted F-V curves were calculated as F(V)w= (F(V)1×z1 + F(V)2×z2 + F(V)3×z3 + F(V)4×z4) / (z1 + z2 + z3 + z4), where F(V)i represents the F(V) function of the ith domain and z; the valence of F(V)i, determined from the Boltzmann-fit y = 1 / (1 + exp[-z×(V-V1/2)/kT]), where V1/2 is the half-activation voltage and k is the Boltzmann-constant and T is temperature.

Statistical significance was attributed to p<.05 as determined by a two-tailed Student’s t-test. For the comparison of multiple groups one-way ANOVA was applied and in the case of a significant difference it was followed by post-hoc Holm-Sidak pairwise comparisons among all groups or versus control. Errors in text and error bars in figures represent 95% confidence interval with number of trials [n] in brackets.

References
Figure S1. G-V and SSI functions of the unlabeled fluorescence constructs

G-V and SSI functions were obtained using the protocols described in the text from unlabeled LFS constructs (circles). Points represent mean ± 95CI from groups of 3 to 6 cells. The dotted line shows the WT-LFS data, while the dashed lines are curves from TAMRA-MTS labeled constructs. For DIV-LFS the SSI curve did not saturate at negative potentials (filled circles), but after DTT treatment (empty circles) approached the labeled channel.
Figure S2. Comparison of integrated gating charge movement for WT-LFS and fluorescence constructs.

Voltage-dependence of integrated gating charge movement for WT-LFS, DI-LFS, DII-LFS, DIII-LFS and DIV-LFS channels. Gating currents were recorded during 20-ms depolarizing steps ranging from -150 to +50 mV in 20-mV steps. Capacitance and leak were removed using P/4 leak subtraction with a sub-sweep holding level of +40 mV. Gating currents were integrated for 10 ms following the voltage change. See Table 1 in manuscript for parameters. Each point represents the mean ± 95CI from 4 to 7 cells.
Figure S3. Comparison of ionic current and fluorescence kinetics in the four domains.

$\text{Na}^+$ currents and fluorescence signals were recorded from DI-LFS, DII-LFS, DIII-LFS and DIV-LFS channels to compare the kinetics of activation. Traces were recorded during a voltage step to +30 mV from -120 mV. The first 10 ms duration is shown.
Figure S4. Comparison of ionic current activation kinetics of DII-LFS and the BrS mutants.

Activation kinetics were measured as $t_{10-90\%}$ rise time following a step to the indicated potentials from a prepulse of -120 mV for DII-LFS, DII-LFS-A735V and DII-LFS-G752R. Each point represents the mean ± 95CI from groups of 3 to 7 cells.
Figure S5: Steady-state inactivation, activation, and rise times for BrS mutants compared to wild-type expressed in HEK293T cells.

(A) Activation kinetics were measured as t\textsubscript{10-90\%} rise time following a step to the indicated potentials from a prepulse of -120 mV for WT [n=4], A735V [n=3], and G752R [n=4].

(B) Voltage-dependence of steady-state activation and steady-state inactivation curves for the WT, A735V, and G752R (purple, green, and red respectively). G-V curves were constructed from I-V relationships recorded from cells depolarized in 10 or 20 mV increments from a prepulse potential of -120 mV. Na\textsuperscript{+} reversal potential was determined for each cell individually. For the SSI curves, cells were held at the conditioning potential ranging from -120 to +20 mV for 50 ms and the available channel fraction was assessed by a depolarizing pulse to -20 mV. Each point represents the mean ± 95CI from 4 to 11 cells.