Structural Mechanism for the Regulation of HCN Ion Channels by the Accessory Protein TRIP8b

Graphical Abstract

Highlights

- DEER and EPR show TRIP8b binds near the C helix on the CNBD of HCN channels
- NMR shows TRIP8b binding affects the C helix and the cAMP binding site
- TRIP8b likely alters both cAMP binding and CNBD conformational changes

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In Brief

Binding of TRIP8b reduces the cyclic nucleotide dependence of hyperpolarization-activated cyclic nucleotide-gated channels. DeBerg et al. identify the interaction topology and suggest that TRIP8b regulates these channels by disrupting both cAMP binding and the coupling of cAMP binding to channel opening.
Structural Mechanism for the Regulation of HCN Ion Channels by the Accessory Protein TRIP8b

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http://dx.doi.org/10.1016/j.str.2015.02.007

SUMMARY

Hyperpolarization-activated cyclic nucleotide-gated (HCN) ion channels underlie the cationic $I_h$ current present in many neurons. The direct binding of cyclic AMP to HCN channels increases the rate and extent of channel opening and results in a depolarizing shift in the voltage dependence of activation. TRIP8b is an accessory protein that regulates the cell surface expression and dendritic localization of HCN channels and reduces the cyclic nucleotide dependence of these channels. Here, we use electron paramagnetic resonance (EPR) to show that TRIP8b binds to the apo state of the cyclic nucleotide binding domain (CNBD) of HCN2 channels without changing the overall domain structure. With EPR and nuclear magnetic resonance, we locate TRIP8b relative to the HCN channel and identify the binding interface on the CNBD. These data provide a structural framework for understanding how TRIP8b regulates the cyclic nucleotide dependence of HCN channels.

INTRODUCTION

In the brain, hyperpolarization-activated cyclic nucleotide-gated (HCN) ion channels contribute to rhythmic firing, dendritic excitability, and resting membrane potential in many different types of neurons (Robinson and Siegelbaum, 2003). These cellular functions play critical roles in learning, sleep, and depression (He et al., 2014; Lörrincz et al., 2002; Nolan et al., 2003). The expression and regulation of HCN channels varies dramatically among different neuronal subtypes. In pyramidal and neocortical neurons, they show a large gradient of expression that can be as much as 60-fold higher in the distal dendrites than in the soma (Lörrincz et al., 2002). In many pathological conditions, including epilepsy and Parkinson’s disease, disruption of the gating, regulation, or localization of HCN channels plays a crucial role in neuronal dysfunction (Chan et al., 2011; He et al., 2014; Lörrincz et al., 2002; Nolan et al., 2003; Reid et al., 2012).

Two features differentiate HCN channels from other members of the voltage-activated channel family: (1) they open in response to membrane hyperpolarization as opposed to depolarization, and (2) they are allosterically regulated by the direct binding of 3’,5’-cyclic-AMP (cAMP) (Craven and Zagotta, 2006; Robinson and Siegelbaum, 2003). Binding of cAMP to HCN channels increases the rate and extent of activation and shifts the voltage dependence of activation to more depolarized voltages (Craven and Zagotta, 2006). The cyclic nucleotide dependence of HCN channels arises from a series of conformational changes in a pair of intracellular domains known as the cyclic nucleotide binding domain (CNBD) and the C linker (Figure 1A). The C linker connects the CNBD to the channel pore and is critical for coupling the conformational changes associated with binding of ligand in the CNBD to the opening of the pore (Craven et al., 2008; Puljung and Zagotta, 2013; Puljung et al., 2014; Taraska and Zagotta, 2007). The CNBD and C linker are characteristic features of the cyclic nucleotide-regulated family of channels, which consists of HCN, cyclic nucleotide-gated (CNG), and KCNH channels (Craven and Zagotta, 2006).

In addition to their activation by membrane hyperpolarization and cAMP, HCN channels are also regulated by a cytoplasmic accessory protein called TRIP8b (tetratricopeptide repeat-containing Rab8b-interacting protein) (Santoro et al., 2004). TRIP8b is a highly alternatively spliced protein that regulates HCN channel cell surface expression in an isoform-dependent manner (Lewis et al., 2009; Santoro et al., 2009; Zolles et al., 2009). TRIP8b has been shown to co-localize with HCN1 channels in the distal dendrites of hippocampal pyramidal neurons, and TRIP8b knockout mice show a striking disruption in this expression pattern (Lewis et al., 2011; Piskorowski et al., 2011).

Structurally, TRIP8b is known to interact at two sites on the carboxy-terminal region of HCN channels (Bankston et al., 2012; Han et al., 2011; Lewis et al., 2009; Santoro et al., 2011) (Figure 1A). A series of tetratricopeptide repeats (TPRs) on TRIP8b interact with the terminal tripeptide sequence of HCN channels with high affinity. This site is thought to anchor TRIP8b onto HCN channels. The structure of this C-terminal interaction has been determined at atomic resolution (Bankston et al., 2012). The second site involves an interaction between a segment of TRIP8b containing residues 223–303, termed TRIP8bcore (Figure 1A and Figure S1A), and the CNBD. This interaction dramatically reduces the cAMP regulation of HCN channels (Lewis et al., 2009; Santoro et al., 2009; Zolles et al., 2009). Biochemical data have suggested that TRIP8b reduces cAMP binding by direct competition with the cAMP binding
site (Han et al., 2011), whereas electrophysiological evidence suggests that TRIP8b alters the effect of cAMP by inhibiting the conformational change in the CNBD associated with cAMP binding (Hu et al., 2013; Zolles et al., 2009). A nuclear magnetic resonance (NMR) study by Saponaro et al. (2014) suggests that TRIP8b does not interact with the cAMP binding site of HCN2 and therefore does not directly compete with cAMP for binding. However, the structure of the CNBD-TRIP8b complex and the mechanism of how TRIP8b regulates cAMP dependence are currently unknown.

In this study, we determine the structural details of the CNBD-TRIP8b complex. We use continuous-wave electron paramagnetic resonance (CW EPR) and double electron-electron resonance (DEER) spectroscopy to examine the conformational transition in the CNBD of HCN2 channels in response to cAMP and TRIP8b binding. DEER is a pulse EPR technique capable of measuring absolute distances and distance distributions between two paramagnetic spin labels attached to a protein and separated by 15–80 Å (Martin et al., 1998; Milov et al., 1984; Pannier et al., 2000). Among distance measurement techniques, a strength of DEER is that it reports a distribution of distances, not merely an average distance. Information on conformational heterogeneity not accessible in crystal structures is fully captured and resolved in DEER distributions from frozen solutions. From DEER measurements between two residues on HCN2, we show that TRIP8b binds to the CNBD in the apo state without significantly altering the overall structure of the domain. Using inter-protein DEER measurements between the CNBD and TRIP8b, we spatially localize two TRIP8b positions relative to the CNBD. Using NMR, we identify CNBD residues involved in TRIP8b binding. These data reveal a binding interface on the CNBD and the topology of the CNBD-TRIP8b complex. Together, they provide mechanistic insights into how TRIP8b causes a decrease in the cAMP dependence of HCN channels.

RESULTS

TRIP8bcore Modulates HCN2 Channels

The TRIP8bcore fragment has been shown to be sufficient to bind to the CNBD of HCN2 channels and antagonize the cyclic nucleotide dependence of these channels (Hu et al., 2013). It contains a central portion of TRIP8b that is highly conserved across species (Figure S1A). To demonstrate that human TRIP8bcore purified from Escherichia coli can bind to and regulate full-length HCN2 channels, we directly applied TRIP8bcore to inside-out patches excised from Xenopus laevis oocytes expressing HCN2 channels. Figure 1B shows HCN2 currents in response to a series of voltage steps to potentials between −70 and −140 mV. In control solutions, HCN2 channels opened slowly with a half-activation voltage of −130 ± 2 mV (Figure 1C). Simultaneous perfusion with 10 μM TRIP8bcore and 1 μM cAMP resulted in an almost complete inhibition of the effects of cAMP, shifting the voltage dependence nearly back to control levels (−127 ± 2 mV, Figure 1C). These data are consistent with previous reports and show that this small fragment of TRIP8b is sufficient to bind to and regulate full-length HCN channels.

Intra-CNBD DEER Probes TRIP8b Binding to the CNBD

The CNBD is composed of an eight-stranded β roll, followed by two α helices (Figure 2A) (Zagotta et al., 2003). Initial ligand binding is thought to take place at the phosphate binding cassette (PBC), which is composed of the region after β6, the P helix, and a small part of α7 (Figure S1B). After ligand binding the last helix in the domain, the C helix, moves closer to the β roll. This motion is thought to be coupled to channel opening (Flynn et al., 2007; Zhou and Siegelbaum, 2007).

Figure 1. TRIP8bcore Binds to and Regulates HCN2 Channels

(A) Cartoon showing the interactions between HCN channels and full-length TRIP8b. The intracellular C linker and the CNBD of HCN2 are labeled. Only two of the four subunits are shown. The variable, core, and TPR domains of TRIP8b are highlighted (see also Figure S1).

(B) Representative current traces elicited by hyperpolarizing voltage steps from inside-out patches of oocytes expressing HCN channels with no ligand (left), in the presence of 1 μM cAMP (center), and in the presence of both 1 μM cAMP and 10 μM TRIP8bcore (right).

(C) Normalized conductance-voltage relationships of HCN alone (cyan), in the presence of 1 μM cAMP (red), and in the presence of both 1 μM cAMP and 10 μM TRIP8bcore (black). Data are represented as means ± SEM.
We have previously shown that a cysteine-free version of the HCN2 CNBD and C linker is able to bind cAMP and undergo conformational change (Flynn et al., 2007; Puljung and Zagotta, 2013; Puljung et al., 2014). To examine whether conformational changes in HCN2 channels accompany TRIP8b binding, we performed DEER experiments on a fragment of HCN2 channels, termed HCN2-CNBDxt, which we expressed and purified from bacteria. This fragment contains the CNBD and helices C through F (residues 488–640, Figure S1B).

Compared with the larger cytosolic fragment (CNBD + C linker), HCN2-CNBDxt has the advantage that it lacks the A and B helices required for tetramerization and that it exhibits enhanced solubility when bound to TRIP8bcore. These advantages are required for our EPR and NMR experiments. Tetramerization would lead to multi-spin effects that complicate EPR results. Also, tetramers would be too large for NMR experiments.

We engineered pairs of cysteines into a cysteine-free version of HCN2-CNBDxt and labeled them with the nitroxide spin label MTSLS ($S\cdot(1\cdotoxy\cdot2,2,5,5\cdottetramethyl\cdot2,5\cdotdihydro\cdot1\cdotH\cdotpyrrol\cdot3\cdoty)(methyl\ methane\ sulfonothioate)$. Cysteines were placed at either of two positions on the C helix (A624C or R635C) and at one of three positions on the β roll (V537C, S563C, or K570C). Figure 2B shows the positions of a single pair of cysteines (V537C and R635C) with spin labels attached. For each spin-labeled site, side-chain ensembles were predicted using the MMM rotamer library (Polyhach et al., 2011). These surface sites were selected based on MMM predictions of favorable labeling and to provide good coverage of the β roll and C helix. The predicted rotamers are shown with spheres indicating the midpoints of the N-O bond on the spin label, corresponding to the positions of the unpaired electron. DEER measures the separation of the unpaired electrons on the N-O group. For DEER experiments, protein samples are snap-frozen, capturing the distribution of backbone and side-chain conformations present at room temperature.

To verify that our MTSLS-labeled HCN2-CNBDxt constructs still bind TRIP8b, we performed fluorescence size-exclusion chromatography on wild-type, V537C/R635C, and S563C/A624C HCN2-CNBDxt. HCN2-CNBDxt and TRIP8bcore were mixed together and passed through a Superdex 200 column with an inline fluorescence detector set to detect tryptophan fluorescence (excitation 280 nm, emission 350 nm). HCN2-CNBDxt contains no tryptophan residues while TRIP8bcore contains five tryptophans. Each chromatogram shows TRIP8bcore alone, wild-type HCN2-CNBDxt alone, and the mixture of the two proteins together (Figure S2). TRIP8bcore elutes at 18.3 ml
when run alone and 16.6 ml when run in combination with WT HCN2-CNBDxt, indicating a stable complex (Figure S2A). Similar results were obtained for the other MTSL-labeled proteins tested (Figures S2B and S2C).

To test the binding of TRIP8bcore compared with full-length TRIP8b(1a-4), we performed DEER experiments on HCN2-CNBDxt labeled at positions 537 and 635, in the presence of 40 μM full-length TRIP8b(1a-4) or 200 μM TRIP8bcore. Modulation-depth normalized DEER time traces for apo- (cyan) and TRIP8b-bound HCN2-CNBDxt are shown in Figure 2C for TRIP8b(1a-4) and Figure 2D for TRIP8bcore. The unnormalized time traces are shown in Figures S3A and S3B. In each case, binding of TRIP8b caused a slower oscillatory component of the time trace to emerge. As the oscillation frequency is proportional to the inverse cube of the distance, these data indicate that a fraction of the spin labels at these two positions moved further apart. DEER distance distributions calculated from the time traces have a maximum at 37 Å in the apo configuration, but are bimodal in the presence of TRIP8b(1a-4) (36 and 46 Å) or TRIP8bcore (36 and 44 Å) (Figures 2E and 2F). A peak near 55 Å can be observed in the distance distribution for TRIP8bcore. However, the uncertainty associated with this peak is large, as indicated by the shaded band. Therefore, this peak is not significant. The similarity of the effect of TRIP8b(1a-4) and TRIP8bcore binding suggests that both proteins bind HCN2-CNBDxt in a similar manner. We performed the remainder of the experiments using TRIP8bcore, which expresses in substantially larger quantities than TRIP8b(1a-4) and contains no native cysteines.

DEER measurements were made for the remaining pairs of positions on HCN2-CNBDxt in the apo, cAMP-bound, and TRIP8bcore-bound states. First, upon cAMP addition, residue 624 at the proximal C helix moved 9 Å closer to the β roll, and residue 635 at the distal C helix made a smaller 1-Å movement toward the β roll (Figures 3A–3C and Figures S3C, S3D, and S3E red traces). This is consistent with similar results obtained with an HCN2 channel fragment consisting of the entire C linker and CNBD (Puljung et al., 2014). No significant cAMP-induced distance changes were seen between a pair of labels on the β roll (Figure 3D and Figure S3F, red traces). These data suggest that the smaller HCN2-CNBDxt fragment behaves similarly to the HCN fragment that contains the full C linker and CNBD.

Addition of TRIP8bcore to HCN2-CNBDxt V537C/A624C and S563C/A624C produced distance distributions that more closely resemble the apo rather than the cAMP-bound distributions (Figures 3B and 3C, black traces). Similar to cAMP, TRIP8bcore binding to HCN2-CNBDxt had no impact on the distance distributions between 537 and 570 on the β roll (Figure 3D, black trace).

Interestingly, for the separation between residue 635 (distal C helix) and residue 563 (β roll), the addition of TRIP8bcore caused a significant sharpening of the distance distribution accompanied by a small shift in the distribution maximum (30 to 33 Å, Figure 3A). In both HCN2-CNBDxt mutants spin labeled at residue
The R635C spectrum is seen upon addition of TRIP8bcore in the presence and absence of 200 nM TRIP8bcore. CW EPR spectra for spin labels attached to indicated residues on HCN in the absence (cyan) and presence (black) of TRIP8bcore. Significant line broadening of the R635C spectrum is seen upon addition of TRIP8bcore.

To determine whether TRIP8b binding affects the mobility of a spin label at the distal end of the C helix, CW EPR spectra were obtained for spin labels attached to indicated residues on HCN in the presence and absence of 200 nM TRIP8bcore. Significant line broadening of the R635C spectrum is seen upon addition of TRIP8bcore.

TRIP8bcore Binds near CNBD Residue 635

To determine whether TRIP8b binding affects the mobility of the spin label at 635, we measured the room temperature CW EPR spectrum of each spin-labeled position of HCN2-CNBDxt in the presence and absence of 200 nM TRIP8bcore. The addition of TRIP8bcore to HCN2-CNBDxt did not result in observable changes in spin-label mobility at any sites except R635C (Figure 4). The R635C spectrum, however, showed significant broadening upon addition of TRIP8bcore, indicating a reduction in MTSL mobility at that position. TRIP8bcore addition increased the rotational correlation time of the spin label from 0.8 to 3.0 ns. This spectral broadening could result from a change in the mobility of the distal C helix or from a significant change in the environment around the spin label at R635C that alters the rotameric distribution of MTSL. In either case, the data suggest that TRIP8bcore binds near R635C on the distal C helix.

Inter-Protein DEER Measurements Localize the Position of TRIP8bcore Bound to HCN2-CNBDxt

To determine the geometry of the HCN2-CNBDxt-TRIP8b complex, we measured DEER distributions between singly spin-labeled HCN2-CNBDxt and singly spin-labeled TRIP8bcore. Seventeen residues on HCN, shown in Figure 5A, and two residues on TRIP8bcore (A248C and A261C) were selected. Both TRIP8bcore variants bound to each labeled HCN2-CNBDxt protein and generated DEER oscillations (Figure S5), with the exception of TRIP8bcore A248C and HCN2-CNBDxt A624C, which showed no oscillations when the two proteins were combined. The resulting DEER distance distributions (Figure 5B) feature main peaks at distances between 20 and 45 Å and suggest that TRIP8bcore is localized near the C helix.

Using the set of inter-protein DEER data, we proceeded to determine more quantitatively the position of TRIP8b relative to HCN2-CNBDxt. We developed a novel probabilistic trilateration algorithm for locating the spin label on a protein relative to a binding partner with a known structure. We generated three-dimensional spatial probability distributions for spin labels attached to TRIP8bcore by combining the DEER distributions with the known structure of the HCN2 CNBD. Briefly, rotamer-weighted radial probability distributions were created from the DEER data for every pair of sites. We based our HCN spin label positions on a model of the apo state of the CNBD determined from previous DEER experiments (Puljung et al., 2014). For each TRIP8bcore site, an overall spatial probability density function was then calculated by multiplying the individual radial probability distributions. Since we do not have accurate structural information about the distal C helix owing to its disorder in the apo state, we excluded 635 from our calculation.

Using this probabilistic method, we located the spin labels attached to TRIP8bcore, A248C, and A261C, relative to HCN2-CNBDxt. Isosurfaces representing regions of constant probability density where the spin labels are likely to be found are shown in Figure 6. The label attached to TRIP8bcore A248C resides near the proximal portion of the C helix near residue A624 on HCN2-CNBDxt (Figure 6, yellow surface). This result is consistent with the possibility that this labeled construct did not bind to the MTSL-labeled HCN2-CNBDxt A624C due to steric hindrance. Position A261C is localized near the β4-β5 loop and the distal end of the C helix (Figure 6, green surface). A binding site near the distal C helix is consistent with our CW EPR observation of restricted spin-label mobility at residue 635 of HCN2-CNBDxt in the presence of TRIP8bcore. A decrease in spin-label mobility upon TRIP8bcore binding would also account for the changes we observed in the DEER distribution between residue 635 of HCN2-CNBDxt and residues on the β roll of HCN2-CNBDxt. Recalculating the isosurfaces using a recent solution state NMR structure of the apo HCN2-CNBD also showed similar probability densities for these residues on TRIP8bcore along the C helix (data not shown). In addition, including the data from the R635C distributions did not dramatically alter the predicted surfaces. We calculated expected distance distributions from the isosurface-bound volumes in Figure 6 and compared them with our experimental distance distributions (Figure S6). The good agreement confirmed the robustness of our probabilistic trilateration analysis.

NMR Identifies a TRIP8bcore Binding Surface on the CNBD

To further examine the interface between the core domain of TRIP8b and the CNBD of HCN channels, we used NMR...
spectroscopy to map the TRIP8b binding site on HCN4-CNBDxt, a similar human HCN4 CNBD construct and the closely related human HCN2 CNBD have recently been published (Akimoto et al., 2014; Saponaro et al., 2014). We confirmed backbone assignments for the HCN4-CNBDxt under our experimental conditions with a transverse relaxation-optimized spectroscopy (TROSY)-HNCA experiment recorded on $^{15}$N,$^{13}$C-labeled HCN4-CNBDxt. We were able to obtain assignments for most (79%) of the HCN4 backbone resonances. Though observed, we were unable to confirm assignment of the disordered N- and C-terminal ends of the protein construct. We were also unable to confirm assignments previously reported for three residues (660–662) corresponding to the P helix (Saponaro et al., 2014).

We collected a series of $^1$H,$^{15}$N-HSQC (heteronuclear single quantum coherence) spectra of 100 μM $^{15}$N-labeled HCN4-CNBDxt in the presence of increasing concentrations of unlabeled TRIP8b$_{core}$. Significant resonance perturbations, primarily broadenings, were observed at low concentrations of TRIP8b$_{core}$ (0.25 molar equivalents), with substantial loss of signal at 0.5 molar equivalents of TRIP8b$_{core}$ and new resonances beginning to appear at higher concentrations (Figure S6). These observations suggest the interaction between HCN4-CNBDxt and TRIP8b$_{core}$ is in the intermediate to slow exchange regime on the NMR timescale under these experimental conditions. In the intermediate exchange regime, the observed line width for a peak will be the sum of the population-weighted average of the line widths of the free and bound species and an additional chemical exchange contribution that scales as $f_f \Delta \omega_{dx}^2$, where $f_f$ and $f_b$ refer to the mole fractions of the free and bound species, and $\Delta \omega_{dx}$ is the difference in chemical shift, in
TRIP8b has been shown to reduce the cAMP dependence of HCN channels. Interactions between a small central region of TRIP8b and the CNBD of HCN are critical for this gating effect (Hu et al., 2013). The location of the binding site for TRIP8b on the CNBD and the mechanism through which TRIP8b exerts its regulatory effect on HCN channels are still not resolved. To answer these questions, we examined how TRIP8b binding affected the overall structure of the CNBD using intramolecular DEER on HCN2-CNBDxt. In addition, using inter-protein DEER on TRIP8bcore and HCN2-CNBDxt, we localized two positions on TRIP8b relative to the CNBD. Lastly, using NMR we found a set of CNBD residues that are at the binding surface for TRIP8b.

The location of the binding surface suggests interesting mechanistic possibilities for how TRIP8b reduces the cAMP dependence of these channels. A model for activation of HCN channels by cAMP has been developed using structural, biochemical, and electrophysiological data (Akimoto et al., 2014; Flynn et al., 2007; Matulef et al., 1999; Saponaro et al., 2014; Tibbs et al., 1998; Zhou and Siegelbaum, 2007). In this model, cAMP initially interacts with residues on the β roll and P helix that compose the PBC. Mutations in this region, including residues in β5 and the PBC, have been shown to reduce cyclic nucleotide affinity (Matulef et al., 1999; Tibbs et al., 1998; Xu et al., 2010; Zhou and Siegelbaum, 2007). After initial ligand binding, the C helix is believed to move toward the binding pocket to form more contacts with the ligand. Mutations in the C helix have been shown to alter ligand affinity and affect the ability of the ligand to promote channel opening (Matulef et al., 1999; Zhou and Siegelbaum, 2007).

Previous work has proposed two mechanisms for how TRIP8b could reduce cAMP dependence of HCN channels. Han et al. (2011) proposed that TRIP8b could be competing with cAMP for the binding site, thus directly reducing its affinity. Hu et al. (2013) and Saponaro et al. (2014) have suggested that TRIP8b allosterically reduces the effect of cAMP by stabilizing the apo conformation of the HCN channel CNBD, possibly by restricting the motion of the C helix. Our results suggest that TRIP8b could be both altering the binding site for the ligand and reducing the ability of the CNBD to undergo the conformational transition associated with channel activation. These data could explain why the physiological effects of TRIP8b could not be fit by a simple allosteric model but required a TRIP8b-specific effect on cAMP binding (Hu et al., 2013). NMR titrations show that the resonances most perturbed by addition of TRIP8b map to the C helix, the PBC, and the β4-β5 loop of HCN4-CNBDxt (Figure 7). This is in excellent agreement with the model generated from the DEER and EPR data, which localizes residue 248 of TRIP8b near the proximal C helix and residue 261 near the distal C helix and the β4-β5 loop. This binding interface and the trilateration-based interaction topology are compatible with the known 4:4 HCN/TRIP8b stoichiometry in the tetrameric HCN channel (Bankston et al., 2012) (Figure 8).

The specific NMR resonances on the CNBD of HCN4 altered by TRIP8b binding include amino acids that have been previously shown to affect both the binding of cAMP and the conformational changes associated with channel gating. Many residues across the C helix, a region that undergoes conformational changes associated with cAMP binding, are affected. For example, an isoleucine in the C helix (636 in HCN2-CNBDxt...
and 714 in HCN4-CNBDxt) that is important for ligand selectivity and affinity is affected by TRIP8b binding (Flynn et al., 2007; Zagotta et al., 2003; Zhou and Siegelbaum, 2007). In addition, an arginine (632 in HCN2-CNBDxt and 710 in HCN4-CNBDxt) that has been shown to form contacts with the ligand and the core of the β roll and to stabilize the open conformation of the channel is affected (Zhou and Siegelbaum, 2007). These results are similar to previous data from the homologous CNBD of human HCN2 (Saponaro et al., 2014), which also show extensive changes in C helix resonances upon addition of TRIP8b. However, our data suggest that TRIP8b binding also affects CNBD regions involved in initial cAMP binding. Specifically, we identify resonance changes in three residues in the PBC and three more in the β4-β5 loop. Each of these regions contains residues that form direct contacts with the cyclic nucleotide (Xu et al., 2010; Zagotta et al., 2003). By affecting both the binding site for cAMP and the C helix, TRIP8b could reduce the cyclic nucleotide dependence of HCN channels via two mechanisms: it could disrupt ligand affinity and interfere with the ability of the C helix to couple ligand binding to channel opening.

The binding interface between the CNBD and TRIP8b points to a possible role of electrostatics in the interaction. TRIP8b_core is highly negatively charged (predicted pI 4.88), with three negatively charged residues near position 261 (EWEE) and another four negatively charged residues between position 248 and 261. The distal portion of the C helix contains a number of positive charges, including two arginines (632 and 635 in HCN2-CNBDxt and 710 and 713 in HCN4-CNBDxt) whose resonances changed in our NMR experiments. Previous results show that charge-inverting mutation of two lysines after the C helix to glutamates (positions equivalent to 638 and 639 in HCN2-CNBDxt and 716 and 717 in HCN4-CNBDxt) resulted in significantly decreased TRIP8b_core binding (Saponaro et al., 2014). In addition, the β4-β5 loop contains two positively charged lysines that are also perturbed in our NMR data. The cAMP binding pocket is highly positively charged and is likely necessary for the binding of the negatively charged cAMP. It is possible that the negatively charged TRIP8b could also lower cAMP affinity by altering the electropositive nature of the binding pocket.

The DEER trilateration and NMR perturbation results in this study provide new data on the interaction topology of the complex formed between the CNBD and TRIP8b. In addition, the intra-CNBD DEER data show that TRIP8b binds to a conformation that is largely similar to the apo state. The binding interface is composed of regions on the CNBD that are critical for ligand binding and regions critical for the ability of the CNBD to couple binding to channel activation. These data provide a structural basis for understanding how TRIP8b interacts with HCN channels. They suggest that TRIP8b might be altering the cAMP dependence of HCN channels by both restricting ligand binding and interfering with the conformational transition necessary to couple cAMP binding to channel activation.

**EXPERIMENTAL PROCEDURES**

**Molecular Biology**

All constructs used in this study were subcloned into the pETM11 vector. The proteins were separated from the histidine tag by a tobacco etch virus (TEV) cleavable linker. TRIP8b_core was derived from the TRIP8b(1a-4) isoform and contains residues 223–303 of that protein. Murine HCN2-CNBDxt contains residues 488–640 in a cysteine-free background (Taraska et al., 2009). Human HCN4-CNBDxt is composed of wild-type residues 563–727. Cysteine mutations were engineered into the constructs at the indicated positions using standard PCR-based methods. All constructs were confirmed with...
fluorescence-based automated sequencing. The cDNA encoding the full-length murine HCN2 channel in the pHE vectors were kindly provided by Steven Siegelbaum and Bina Santoro (Columbia University).

Electrophysiology
The cRNA for HCN2 channels was transcribed using the mMessage Machine T7 transcription kit (Ambion) and expressed in X. laevis oocytes that were defolliculated and injected with the cRNA as previously described (Zagotta et al., 1989). The vitelline membranes were manually removed, and currents were measured in cAMP following steps up to 140 mV. A Boltzmann function, 

\[ \frac{G}{G_{\text{max}}} = \frac{1}{1 + \exp \left( \frac{V - V_{1/2}}{k} \right) } \]

was then fitted to the data, where \( V \) is the test voltage, \( V_{1/2} \) is the half-maximal activation voltage, and \( k \) is the slope factor.

Protein Expression, Purification, and Spin Labeling
For each protein expressed, the construct was transfected into BL21 (DE3) cells and grown at 37°C to an optical density of 0.6–0.8. The cells were then induced with 1 mM isopropyl \( \beta \)-D-1-thiogalactopyranoside and grown over-night at 18°C. For NMR experiments, the bacteria were grown in MOPS minimal media supplemented with \( ^{15} \text{NH}_4\text{Cl} \) (1 g/l) and \( ^{13} \text{C}_6\)-glucose (2 g/l). After growth and expression, 1- to 2-l cultures of cells were pelleted by centrifugation at 4,000 \( \times \) \( g \) at 4°C for 10 min and resuspended. For EPR experiments, cells were resuspended in 150 mM KCl and 30 mM HEPES at pH 7.4 (for HCN2) or pH 8.5 (for TRIP8bcore). DNase at a final concentration of 5 \( \mu \)g/ml and two tablets of protease inhibitors (cComplete EDTA-free; Roche) were added to the buffer. The resuspended cells were lysed by an Emulsiflex-C3 homogenizer (Avestin) and clarified by centrifugation at 186,000 \( \times \) \( g \) at 4°C for 45 min. The lysate was then purified on an Ni\(^2+\)-affinity resin column (HisTrap HP, GE Healthcare). The octahistidine tag was removed by TEV protease cleavage overnight at 4°C. The protein (10–50 \( \mu \)M) was then spin labeled with 100 \( \mu \)M MTSL (Toronto Research Chemicals) per cysteine mutation for 1 h at room temperature or 6–16 h at 4°C. To remove the TEV protease and further purify the samples, the proteins were passed through ion exchange columns. For HCN2 and HCN4 proteins, a cation exchange column was used (HiTrap SP FF, GE Healthcare). The protein was loaded into 1-mm o.d. quartz capillaries (Sutter, Q165-115-10) and flash-frozen in liquid nitrogen.

EPR Sample Preparation
For CW EPR and DEER experiments, the protein was buffer exchanged into D\(_2\)O with 150 mM KCl, 30 mM Tris, and 10% glycerol (pH 8.4) using a PD-10 column (GE Healthcare). Doubly spin-labeled HCN protein for intra-protein DEER measurements was diluted to approximately 50 \( \mu \)M cAMP, 200 \( \mu \)M TRIP8bcore, or 40 \( \mu \)M full-length TRIP8b was added as indicated in the text. For inter-protein DEER measurements, 150 \( \mu \)M singly spin-labeled HCN was combined with 40 \( \mu \)M singly spin-labeled TRIP8bcore. For DEER, 50 \( \mu \)L of each protein sample was inserted into a 1.65-mm outer-diameter (o.d.) quartz tube (Sutter, Q165-115-10) and flash-frozen in liquid nitrogen. For CW EPR, protein was loaded into 1-mm o.d. quartz capillaries (Sutter, Q100-50-7.5) at concentrations of 25 \( \mu \)M HCN2-CNBDxt and 200 \( \mu \)M TRIP8bcore immediately prior to the experiment.

EPR Data Acquisition
CW EPR spectra were recorded on a 9–10 GHz Bruker EMX spectrometer with a dielectric resonator (Bruker ER4123D, 9.78 GHz, Q-factor 2000–4000),
Experiments were performed at room temperature with 0.2 mW incident power, 2 G modulation amplitude, and 100 kHz modulation frequency. DEER data were acquired on a 33–35 GHz Bruker EleXsys E580 spectrometer with an overcoupled dielectric resonator (Bruker EN150TD2, 34.1 GHz, Q-factor 300–700). Experiments were performed at 80 K using a liquid helium cooling system (Oxford). The four-pulse, dead-time free DEER sequence \((\pi/2)_{\text{probe}} - \tau_1 - \pi - \tau_2 - f - \pi_{\text{pump}} - \tau_2 - f - \pi_{\text{probe}} - \tau_2\) was used with 22-ns probe pulses and a 44-ns pump pulse. Pulse delays were 120 ns for \(\tau_1\) and 1800 ns for \(\tau_2\). The delay \(f\) was varied from –60 ns to between 1,800 and 4,000 ns, depending on the experiments, in 10-ns increments. The pump frequency matched the nitroxide spectral maximum. The probe frequency was centered in the resonator dip and was 62 MHz lower than the pump frequency. An eight-step phase cycling protocol combined with extensive averaging at a repetition time of 2 ms was used to collect data. The measurement time for each sample was 10–16 hr.

**Data Analysis**

Rotational correlation times were extracted from CW EPR data via spectral simulation using EasySpin (Stoll and Schweiger, 2006). Probability densities for spin labels attached to TRIPbcore were calculated in MATLAB. Rotameric models of MTSL attached to HCN2-CNBDxt were obtained using MMM (Polyhach et al., 2011). DEER distance distributions were obtained using DeerAnalysis2013 (Jeschke et al., 2006). A homogeneous three-dimensional background was used for background correction. Time traces were converted to distance distributions using Tikhonov regularization, a model-free least-squares approach. The regularization parameter was optimized separately for each data set according to the L-curve criterion. To estimate errors associated with our measurement, the noise in the time domain traces was linearly transformed to the distance domain. The shaded error bars shown in the distance domain correspond to two standard deviations of the time domain noise. Molecular graphics and analyses were performed with UCSF Chimera and PyMOL (Meng et al., 2006; Pettersen et al., 2004; DeLano, 2010).

**Trilateration**

For each TRIPbcore mutation, every DEER distance distribution between HCN2-CNBDxt and TRIPbcore was transformed into a radial probability distribution in three-dimensional space with the origin centered at the MMM predicted N–O bond midpoint of MTSL. For each HCN2-CNBDxt mutation, a weighted average of these distributions was computed using the rotameric populations predicted by MMM as weights. The total probability densities for each TRIPbcore residue were calculated by multiplying the weighted densities computed for each HCN site.

**NMR**

NMR spectra were recorded on a Bruker Avance III spectrometer equipped with a cryoprobe and operating at 600 MHz. All spectra were recorded at 25°C in 30 mM HEPES, 150 mM KCl, and 10% D2O (pH 7.6). The HNCA spectrum was collected on samples containing 300 \(\mu\)M \(^{13}\)C,\(^{15}\)N-labeled HCN4-CNBDxt. HSQC spectra were collected on samples containing a constant concentration (100 \(\mu\)M) of \(^{13}\)C,\(^{15}\)N-labeled HCN4-CNBDxt from the same preparation in the presence of increasing concentrations of TRIPbcore (0, 25, 50, 100 \(\mu\)M). NMR data were processed using NMRPipe/NMRDraw (Delaglio et al., 1995) and visualized using NMRView (Johnson and Blevins, 1994). Resonance perturbation analysis examined changes in peak intensities by determining the ratios of a peak in the TRIPbcore-bound spectrum to its corresponding peak in the free HCN4CNBDxt spectrum. We then scored each peak for its standard Z score using the formula

\[
Z = \frac{X - \mu}{\sigma},
\]

where \(X\) is the peak intensity ratio, \(\mu\) is the mean intensity ratio for all peaks, and \(\sigma\) is the SD of peak intensities. Peaks with Z scores below –1 were considered significantly perturbed.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes six figures and can be found with this article online at http://dx.doi.org/10.1016/j.str.2015.02.007.

**AUTHOR CONTRIBUTIONS**

H.A.D., J.R.B., J.C.R., P.S.B., W.N.Z., and S.S. designed and analyzed the experiments. H.A.D., J.R.B., and J.C.R., carried out the experiments. H.A.D., J.R.B., W.N.Z., and S.S. wrote the manuscript. All authors discussed the results and commented on the manuscript.

**ACKNOWLEDGMENTS**

We thank Stacey Camp for technical assistance, Ellen Hayes for help with the EPR and DEER experiments, and Yoni Haltin for helpful discussions. This work was supported by NIH grants NS074545 (to J.R.B.), EY010329 (to W.N.Z. and S.S.), ST23CA080416-15 (to J.C.R.) R01 GM098501 (to P.S.B.), American Heart Association grants 14IRG1877000 (to W.N.Z.) and 14CSA20380095 (to W.N.Z. and S.S.), the University of Washington (S.S.), and by a fellowship from the Sackler Scholars Program in Integrative Biophysics (to H.A.D.). Chimera is developed by the Resource for Biocomputing, Visualization, and Informatics at the University of California, San Francisco (supported by NIGMS P41-GM103311).

Received: November 14, 2014
Revised: January 13, 2015
Accepted: February 9, 2015
Published: March 19, 2015

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