Mechanism for the inhibition of the cAMP dependence of HCN ion channels by the auxiliary subunit TRIP8b

TRIP8b, an accessory subunit of hyperpolarization-activated cyclic nucleotide-gated (HCN) ion channels, alters both the cell surface expression and cyclic nucleotide dependence of these channels. However, the mechanism by which TRIP8b exerts these dual effects is still poorly understood. In addition to binding to the carboxyl-terminal tripeptide of HCN channels, TRIP8b also binds directly to the cyclic nucleotide-binding domain (CNBD). That interaction, which requires a small central portion of TRIP8b termed TRIP8bcore, is both necessary and sufficient for reducing the cAMP-dependent regulation of HCN channels. Here, using fluorescence anisotropy, we report that TRIP8b binding to the CNBD of HCN2 channels decreases the apparent affinity of cAMP for the CNBD. We explored two possible mechanisms for this inhibition. A noncompetitive mechanism in which TRIP8b inhibits the conformational change of the CNBD associated with cAMP regulation and a competitive mechanism in which TRIP8b and cAMP compete for the same binding site. To test these two mechanisms, we used a combination of fluorescence anisotropy, biolayer interferometry, and double electron-electron resonance spectroscopy. Fitting these models to our fluorescence anisotropy binding data revealed that, surprisingly, the TRIP8b-dependent reduction of cAMP binding to the CNBD can largely be explained by partial competition between TRIP8b and cAMP. On the basis of these findings, we propose that TRIP8b competes with a portion of the cAMP-binding site or distorts the binding site by making interactions with the binding pocket, thus acting predominantly as a competitive antagonist that inhibits the cyclic-nucleotide dependence of HCN channels.

Electrical activity in cells throughout the body requires the precisely regulated opening and closing of ion channels. This tuning of ion channel function can occur in many ways, including the binding of ligands and through the association with accessory proteins. The opening of hyperpolarization-activated cyclic nucleotide-gated (HCN) channels is enhanced by the direct binding of cAMP to a highly conserved cytoplasmic cyclic nucleotide-binding domain (CNBD) in each subunit (1). However, in neurons, this regulation of the channel function is mostly eliminated through the binding of the accessory subunit TRIP8b (2, 3).

TRIP8b is a highly alternatively spliced and primarily neuronal protein that can assemble with all mammalian HCN channels (2–6). In addition to reducing the cAMP dependence of HCN channels, TRIP8b regulates the cell surface expression in an isoform-dependent manner (2, 3, 6). TRIP8b has been shown to co-localize with HCN1 in the distal dendrites of hippocampal pyramidal neurons, and knocking out TRIP8b dramatically disrupts this dendritic localization (4, 5).

TRIP8b has a bipartite interaction with the carboxyl-terminal region of HCN channels. One binding interface for these two proteins involves a series of tetratricopeptide repeats (TPRs) on TRIP8b and the carboxyl-terminal residues of HCN channels (2, 3, 7, 8). The interaction has been elucidated in atomic detail with X-ray crystallography and is thought to be a high-affinity site that anchors these two proteins together (8).

The other binding interface involves the binding of a small segment of TRIP8b that has been termed TRIP8bcore (residues 223–303) to the CNBD of HCN channels (4, 5, 9, 10). Patch clamp experiments utilizing only this TRIP8bcore domain showed that it is both necessary and sufficient to inhibit the cAMP dependence of these channels (9, 10). This interface between the CNBD and TRIP8bcore has been localized using double electron-electron resonance (DEER) and NMR spectroscopy (10, 11). The interface involves regions on the CNBD that are critical for cAMP binding, as well as regions critical for the conformational change that occurs subsequent to cAMP binding.

Although it is known that TRIP8b reduces the cAMP dependence of the channel gating, there is still debate as to the mechanism of this regulation. Biochemical data utilizing pulldown assays showed that increasing the concentration of cAMP or...
mutating the cAMP-binding site both reduced the total TRIP8b pulled down by HCN channels (5). The authors concluded that this suggested a direct competition between TRIP8b and cAMP for binding to HCN channels. In contrast, Hu et al. (9) analyzed patch clamp experiments with a model that suggested a mechanism where TRIP8b does not directly compete with cAMP for the binding to the channel but instead acts through an allosteric mechanism to decrease the affinity of the CNBD for ligand. Recent NMR work from two labs showed similar binding interfaces between the CNBD and TRIP8b, but each study arrived at subtly distinct conclusions (10, 11). Both groups suggested that TRIP8b could be working through both a competitive and non-competitive mechanism. However, the two studies differed in the precise structural hypothesis for this competition. Saponaro et al. (11) suggested that TRIP8b binding might work by inhibiting the conformational change associated with channel opening that takes place in the N-terminal region of the CNBD and allosterically reducing cAMP binding affinity. DeBerg et al. (10) hypothesized a partial overlap of the binding sites for TRIP8b and cAMP and inhibition of a different conformational change in the C-helix that is associated with cAMP binding.

Given this uncertainty, we set out to determine whether TRIP8b acts as a competitive antagonist or noncompetitive antagonist of HCN channels. We used fluorescence anisotropy and biolayer interferometry to quantify the binding affinities of cAMP and TRIP8b for the isolated CNBD of HCN2. In addition, we utilized DEER to quantify the extent to which cAMP and TRIP8b regulate the activation conformational change of the CNBD. With these data, we were able to test the two mechanisms by fitting the data to models for competitive and noncompetitive mechanisms of inhibition. We found that, surprisingly, a partially competitive mechanism, but not a noncompetitive mechanism, was sufficient to explain the TRIP8b inhibition in cAMP binding.

Results
TRIP8bcore inhibits cAMP-dependent regulation of HCN2 channels

TRIP8b is a cytosolic protein composed of three primary domains (Fig. 1A). The first is the variable N-terminal domain that is the site of alternative splicing. The second is a small central domain comprised of 80 amino acids called TRIP8bcore, which contains a conserved region that is present in all orthologs of TRIP8b, as well some flanking sequence on both the amino- and carboxyl-terminal sides of the protein. The last is a carboxyl-terminal series of six TPR repeats that bind to the terminal amino acids at the carboxyl terminus of HCN channels (Fig. 1B). Recent work has shown that TRIP8bcore binds to the CNBD near a critical mobile helix called the C-helix, which changes conformation in response to cAMP binding (Fig. 1B) (10, 11). This binding virtually eliminates the cAMP-dependent regulation of HCN2 channels (9, 10, 12). Here, we confirmed that effect by applying purified TRIP8bcore to inside-out excised patches from Xenopus laevis oocytes expressing HCN2 channels. In the absence of TRIP8b, 1 μM cAMP dramatically increased the rate and extent of channel opening and shifted the voltage dependence of activation to more depolarized levels (Fig. 1C).

However, perfusing both 1 μM cAMP and 10 μM TRIP8b onto the patch resulted in currents that were nearly identical to control currents.

cAMP and TRIP8bcore bind the CNBD

To understand the mechanism of TRIP8b inhibition of the cAMP dependence of HCN channels, we measured the binding affinities of both cAMP and TRIP8b for the CNBD. To do this, we used a fragment of the HCN2 CNBD that contains the CNBD and helices C’ through F’ of the C-linker (residues 488–640) termed HCN2-CNBDxt. We have previously shown that this fragment of HCN2 is able to bind cAMP and undergo the conformational change associated with cyclic nucleotide binding (10). Compared with the larger cytosolic fragment that contains both the CNBD and the full C-linker, HCN2-CNBDxt lacks the region required for domain tetramerization and is more soluble when in complex with TRIP8b.

To determine the apparent affinity of cAMP for the CNBD, we measured the anisotropy of 20 nM 8-fluo-cAMP, a fluorescent analog of cAMP, with increasing amounts of HCN2-CNBDxt. Fluorescence anisotropy was then plotted versus total concentration of HCN2-CNBDxt, and the data were fit with a single binding isotherm (see “Experimental procedures,” Equation 2) (Fig. 2A). This revealed an apparent binding affinity for HCN2-CNBDxt and 8-fluo-cAMP of 324 ± 88 nM (n = 3).

To determine the binding affinity for TRIP8bcore and HCN2-CNBDxt, we labeled TRIP8bcore with the fluorophore bimane. To do this, position Ala-261 of TRIP8bcore was mutated to cys-

\[\text{TRIP8b regulation of HCN channels} \]

\[\text{A Variable domain TRIP8bcore TPR domains} \]

\[\text{B Out} \]

\[\text{C-Linker CNBD TRIP8bcore TRIP8b TPR domains} \]

\[\text{Figure 1. TRIP8b inhibits the cAMP dependence of HCN2 channels.} \]

\[\text{A, schematic cartoon showing the major domains of TRIP8b. The orange rectangle represents the variable domain 1a. The yellow rectangle represents a conserved region that is absolutely conserved in all orthologs of TRIP8b. The light green hexagons represent the individual TPRs that make up the TPR domain. B, cartoon showing TRIP8b interacting with the carboxyl-terminal region of HCN channels. The structure of the C-linker/CNBD is adapted from PDB 1Q43 (30). C, representative current traces elicited by stepping the voltage to hyperpolarized potentials between −70 and −140 mV in inside-out patches from oocytes expressing HCN2 channels. The top panel shows currents in the absence of ligand, the middle panel is in the presence of 1 μM cAMP, and the bottom panel is in the presence of both 1 μM cAMP and 10 μM TRIP8b.} \]
TRIP8b regulation of HCN channels

Figure 2. TRIP8b and cAMP both bind to HCN2-CNBDxt. A, fluorescence anisotropy measurements of a fluorescent analog of cAMP (8-fluo-cAMP) plotted versus the total concentration of HCN2-CNBDxt (n = 3). These data were fit with Equation 2 to give an apparent binding affinity of 324 ± 88 nM. B, fluorescence anisotropy measurements of a bimane-labeled TRIP8b-core plotted versus the total concentration of HCN2-CNBDxt (n = 3). These data were fit with Equation 2 to give an apparent binding affinity of 8.4 ± 4.1 μM. The data are plotted as means ± S.E.

We previously labeled this position with MTS spin labels for EPR experiments and showed that the labeled TRIP8b still bound to the CNBD of HCN channels (10). The anisotropy of the labeled TRIP8b-core (50 nm) was then measured in different amounts of HCN2-CNBDxt. These data were plotted and fit with a single binding isotherm (Equation 2), which showed a binding affinity of 8.4 ± 4.1 μM (n = 3) (Fig. 2B).

As an additional assay for TRIP8b binding, we utilized biosensor interferometry, which analyzes the shift in interference pattern as one protein binds to a fiber optic biosensor prebound with an immobilized second protein. This method allows us to measure not only the affinity of the interaction, but also the rates of binding and unbinding. Another advantage of this approach is that it uses wild-type unlabeled proteins, eliminating the possibility that the fluorescent label could be affecting binding. To measure binding of TRIP8b-core to HCN2-CNBDxt, we immobilized an N-terminally His-tagged HCN2-CNBDxt onto a row of eight biosensors that are coated with Ni-NTA. The biosensors were then dipped into a 96-well plate with wells that contain varying concentrations of TRIP8b-core, which allowed for the measurement of binding (on) rates (Fig. 3A, left panel). The sensor was then moved to a well that contained buffer only and the unbinding (off) rates were determined (Fig. 3A, right panel).

A representative set of binding curves is shown in Fig. 3B. We first used Equation 3 to fit these data. We made k_{on} and k_{off} global fit parameters and determined the values that produced the best fit to all the TRIP8b-core concentrations for a single preparation (Fig. 3C). Doing this with three different preparations of HCN2-CNBDxt and TRIP8b-core resulted in a measured k_{on} of 1 × 10^8 ± 0.16 × 10^8 M^{-1} s^{-1} and k_{off} of 0.72 ± 0.14 s^{-1} (n = 3). From these data, K_d (= k_{off}/k_{on}) for the binding of TRIP8b-core to HCN2-CNBDxt was 7.35 ± 1.3 μM (n = 3), similar to the value measured from anisotropy measurements.

As a second method for calculating the on and off rates, the binding data for each concentration of TRIP8b-core were fit individually with Equation 4. The resulting k_{obs} from the fits were then plotted versus the concentration of TRIP8b-core (Fig. 3D). As expected, k_{obs} was linearly related to protein concentration, and the data could be fit with Equation 5. The slope of this line gives a measurement of k_{on} (1.12 × 10^5 M^{-1} s^{-1}), whereas the y intercept gives k_{off} (0.73 s^{-1}). The K_d value (k_{off}/k_{on}) of 6.5 μM is in good agreement with the global fit calculation, as well as the anisotropy measurement. These data establish the rates of the binding and unbinding of TRIP8b-core to HCN2-CNBDxt and demonstrate that the fluorophore labeling of TRIP8b-core did not significantly affect binding to HCN2-CNBDxt.

TRIP8b inhibits binding of cAMP to HCN2

Functional studies have shown that TRIP8b reduces the apparent affinity of cAMP for activation of HCN channels (9, 10). We sought to determine the mechanism for this inhibition by directly measuring the binding of cAMP to HCN channels in the presence of TRIP8b. Again, we measured the fluorescence anisotropy of 8-fluo-cAMP at increasing concentrations of HCN2-CNBDxt. However, in this experiment, the measurements were made in the presence of increasing amounts of TRIP8b-core, and the data were normalized by subtracting the minimum anisotropy value from each data point and dividing by the maximum change in anisotropy. The results can be seen in Fig. 4A. The apparent affinity of cAMP binding to HCN2-CNBDxt decreases as the total concentration of TRIP8b increases.

To determine the mechanism of this inhibition of cAMP binding, we considered two models for the inhibition: noncom-
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Figure 3. Biolayer interferometry reveals TRIP8bcore affinity for HCN2-CNBDxt. A, cartoon representation of biaxial interferometry experiments. The colored rectangle represents the optical probe, and the green is the layer coated with Ni-NTA. The probes are loaded with HCN2-CNBDxt (blue ovals). The probe is then dipped into a well of a 96-well plate containing TRIP8bcore (yellow rectangles). TRIP8bcore binding kinetics are then measured. The probe is then moved to a well with only buffer, and TRIP8bcore unbinding kinetics are measured. B, binding curves that show the shift in interference pattern of light as a function of time. This shift is directly related to the change in thickness of the optical layer, which is in turn related to the binding of TRIP8bcore. The concentration of TRIP8bcore in the wells is indicated. C, the boxed regions from B are shown at an expanded time scale. These data are fit with Equation 3 revealing the on and off rates of TRIP8b binding. The average rates were $k_{on} = 1 \times 10^5 \pm 0.16 \times 10^5 \text{ M}^{-1} \text{s}^{-1}$ and $k_{off} = 0.72 \pm 0.14 \text{ s}^{-1}$ ($n = 3$). D, plot of $k_{on}$ versus the concentration of TRIP8b ($n = 3$). The slope of the linear fit to Equation 4 gives an estimate of $k_{on}$ of $1.12 \times 10^5 \text{ M}^{-1} \text{s}^{-1}$, and the vertical intercept estimates a $k_{off}$ of $0.73 \text{ s}^{-1}$. These data are consistent with the global fits to the data in B and C. The data are plotted as means $\pm$ S.E.

petitive and partially competitive. In the first model, both cAMP and TRIP8b can bind to HCN2-CNBDxt simultaneously, but TRIP8b inhibits the conformational change associated with cAMP binding, (Fig. 4B). In the second model, TRIP8b and cAMP compete for binding to the CNBD (Fig. 4C). In other words, cAMP inhibits the initial binding of TRIP8b and vice versa, suggesting partial overlap in their binding sites. Fitting these models to the steady-state binding data in Fig. 4A required that we determine the equilibrium constants $K_T$, $K_C$, and $L$. The dissociation constant of TRIP8bcore for the unbound CNBD ($K_C$) was taken from the global fits to the interferometry data (7.3 $\mu$M), but all three of our measurements of TRIP8bcore binding were in such good agreement that the model predictions are unaffected by which value we used. The dissociation constant for cAMP for the resting state of HCN2-CNBDxt ($K_C$) was determined from fitting a three-state model resulting from the models in Fig. 4 (B and C) when [TRIP8bcore] = 0. As described previously, the normalized anisotropy data of 8-fluocAMP binding in the absence of TRIP8bcore were fit using Equation 2 and produced a $K_C$ of 324 nM. This number reflects the apparent affinity of cAMP for HCN2-CNBDxt. To determine $K_C$, we can use this apparent affinity, as well as the equilibrium constant for the activation conformational change, and solve the equation $K_C = K_C^p/(1 + L)$.

To determine $L$, the equilibrium constant for the activation conformational change in the CNBD with bound cAMP, we utilized DEER spectroscopy. DEER measures distance distributions between two spin labels attached to a protein. DEER works over distances between 15 and 80 Å. The distance distributions measured by DEER report multiple conformational states of a protein along with the steady-state probability that the protein adopts each conformation. From these measurements, we can calculate the equilibrium constant between different conformations.

To measure the equilibrium constant for the activation conformational change in the CNBD, we introduced a pair of cysteines into HCN2-CNBDxt and labeled the sites with the spin label S-(1-oxyl-2,2,5,5-tetramethyl-2,5-dihydro-1H-pyrrol-3-yl)methyl methanesulfonothioate (MTSL). For these experiments, we selected a residue in the β-roll (S563C) and a residue in the C-helix (A624C) of the CNBD that have been previously shown to report the conformational change in the CNBD associated with cAMP binding (10, 13, 14). Fig. 5A shows the positions of this pair of cysteines with spin labels attached. For each spin-labeled site, side-chain ensembles were predicted using the MMM rotamer library (15). We have previously shown that when cAMP binds to the CNBD, the C-helix moves closer to the β-roll by ~9 Å (10, 13, 14). Here we performed a similar experiment by looking at the distance distribution between these two positions in the absence and presence of saturating (1 mM) 8-fluocAMP (Fig. 5B). In an attempt to trap the room temperature equilibrium conformational ensemble, samples were flash frozen by rapid placement into liquid nitrogen (16, 17). As with previous experiments, we observed a shift in the distance distribution to shorter distances associated with a conformational change from the resting to the active state. However, even in saturating 8-fluocAMP, a fairly substantial probability density exists in the longer distance conformation. This balance between the longer “resting” state and the shorter “active” state represents an equilibrium between these two conformations that exists even in the presence of saturating ligand.
To quantify this equilibrium, we measured the area under these peaks and calculated $L$ by looking at the ratio of the areas under the activated versus resting components. The distance cutoff for discriminating active versus resting state was set at 31.5 Å. This resulted in a value for $L$ of $1.3 \pm 0.2$ for 8-fluo-cAMP bound HCN2-CNBDxt, similar to the value determined for cAMP (13, 14, 18).

**TRIP8b and cAMP compete for HCN channel binding**

Using the previously determined equilibrium constants, we were then able to test our two models for TRIP8b inhibition of cAMP regulation. In the noncompetitive model in Fig. 6A, we assumed that TRIP8b only altered the resting-to-active conformational change after initial cAMP binding. In other words, the resting-state binding affinity of cAMP without and with TRIP8b core bound was the same ($K_C$), and the resting-state binding affinity of TRIP8b core without and with cAMP was the same ($K_T$). Only the equilibrium constant for the activation conformational change was different, decreased by TRIP8b core binding by a factor of $y$ ($L$ versus $L/y$). Surprisingly, even with a decrease in the equilibrium constant for activation by 4 orders of magnitude when TRIP8b core was bound ($y = 10^4$), the anisotropy data were not well fit by the purely noncompetitive model (Fig. 6A).

Alternatively, we tested the partial competition model by varying the dissociation constant for cAMP binding in the presence of TRIP8b while assuming that the conformational change subsequent to cAMP binding is unaffected. Fig. 6B shows that a 100-fold increase in the dissociation constant for cAMP binding in the presence of TRIP8b core ($x = 100$) is sufficient to fit the data well. A 100-fold effect on the equilibrium constant for binding is not complete competition but represents a 2.7 kcal/mol destabilization of cAMP binding in the presence of TRIP8b. Surprisingly, this suggests that partial competition for the binding site on the channel is sufficient to account for TRIP8b inhibition of the cAMP dependence of HCN channels. It is worth noting that using our models, we were only able to fit the data if TRIP8b core binding changed the cAMP affinity by at least 100-fold, even if it also changed the equilibrium constant for the conformational change. Supplemental Fig. S1 shows an example where we increased the dissociation constant for cAMP binding by 10-fold and decreased the equilibrium constant for the conformational change by 10-fold, and the data were still not fit well. Although this does not eliminate the possibility that TRIP8b could be affecting both the binding of cAMP and the conformational change, our results suggest, in...
the context of our models, that an effect on binding is necessary and sufficient to allow TRIP8b to antagonize the cAMP dependence of HCN channels.

To confirm our conclusion using an alternative approach, we again utilized DEER to measure the probability that the CNBD is in the active conformation. We performed DEER experiments that can distinguish between the two models for TRIP8b inhibition of cAMP regulation. The models make different predictions about the probability that HCN2-CNBDxt would be in the active state. We performed DEER experiments in the absence and presence of TRIP8b and 8-fluo-cAMP. We calculated these probabilities for each model using 25 μM HCN2-CNBDxt and 1 mM cAMP and varied the concentration of TRIP8b. For the noncompetitive model, we assumed $y = 10^4$ and for the partially competitive model, we assumed $x = 10^4$. As shown in Fig. 7A, in the noncompetitive model, increasing the concentration of TRIP8b dramatically reduced the fraction of active CNBD molecules. However, in the partially competitive model, the fraction of CNBD molecules in the active state was insensitive to TRIP8b concentration.

Based on these simulations, we designed DEER experiments to further test which model best explains the mechanism of TRIP8b regulation of HCN channels. We replicated these model conditions in our DEER experiments, using 25 μM HCN2-CNBDxt and 1 mM 8-fluo-cAMP in each experiment. We performed DEER experiments in the absence and presence of 300 μM TRIP8b. At this concentration of TRIP8b, the two models make dramatically different predictions (Fig. 7A). The noncompetitive model predicts that this concentration of TRIP8b should have a large effect on the fraction of CNBD molecules in the active conformation, whereas the partially competitive model predicts that TRIP8b should have little effect. The result of the DEER experiments are shown in Fig. 7B. In this study, we assumed $x = 10^4$ and $y = 10^4$, making the $H_{TCA}$ state essentially unpopulated. The fits are very poor. For the noncompetitive model, $K_C = 0.745 \mu M, K_T = 7.34 \mu M, L = 1.3$, and $y = 10^4$. For the partially competitive model, $K_C = 0.745 \mu M, K_T = 7.34 \mu M, L = 1.3$, and $x = 10^4$. The data are plotted as means ± S.E.

Discussion

TRIP8b has been shown to dramatically reduce the cAMP dependence of HCN channels (2, 3, 5, 8–10, 12). Binding of a small central region of TRIP8b (TRIP8bcore) to the CNBD of HCN channels is both necessary and sufficient for this effect (9, 10). However, the mechanism for this inhibition is still unclear. To better understand the mechanism, we considered two multistate models that represent the simplest possible mechanisms of inhibition. In the first mechanism, noncompetitive inhibition, TRIP8b binds to the CNBD of HCN channels and inhibits the conformational change associated with channel activation. This mechanism is perhaps most intuitive, given that the TRIP8b binding site is thought to involve moving parts of the CNBD (10, 11). In the second mechanism, partially competitive inhibition, TRIP8b and cAMP are competing for the same binding site on the CNBD. Although less intuitive, this mechanism suggests that the binding sites for cAMP and TRIP8b partially overlap or that the binding of one ligand distorts the binding site of the other (through a conformational change different from the one associated with channel activation). Fitting these models to cAMP binding data in the absence and presence of TRIP8bcore, we found that competition was both necessary and sufficient to fit our data.

Beyond suggesting that partial competition explained our binding data, the models we developed also made specific predictions about how the CNBD would change conformation in the presence of cAMP and TRIP8b. If the mechanism was partial competition, the model predicts that, in saturating cAMP, the fraction of CNBD molecules in the active state was insensitive to the concentration of TRIP8b. For the noncompetitive model, $K_C = 0.745 \mu M, K_T = 7.34 \mu M, L = 1.3$, and $x = 10^4$. As shown in Fig. 7A, in the noncompetitive model, increasing the concentration of TRIP8b dramatically reduced the fraction of active CNBD molecules. However, in the partially competitive model, the fraction of CNBD molecules in the active state was insensitive to TRIP8b concentration.

The models make dramatically different predictions (Fig. 7A). The noncompetitive model predicts that this concentration of TRIP8b should have a large effect on the fraction of CNBD molecules in the active conformation, whereas the partially competitive model predicts that TRIP8b should have little effect. The result of the DEER experiments are shown in Fig. 7B. In the presence of 300 μM TRIP8bcore and 1 mM 8-fluo-cAMP, the fraction of HCN2-CNBDxt in the active state is almost identical to that of the domain in the presence of 8-fluo-cAMP alone, consistent with the predictions made by the partially competitive model. These experiments were done using 8-fluo-cAMP to maintain consistency in the experiments throughout this study; however, supplemental Fig. S2 shows that qualitatively the same pattern is seen when using cAMP.

Figure 6. Data for cAMP binding to HCN2-CNBDxt in the presence of TRIP8b are best fit with a partial competition model. A, normalized fluorescence anisotropy data for 8-fluo-cAMP binding to HCN2-CNBDxt at different concentrations of TRIP8bcore fit with the noncompetitive model at the top. In this example, $y = 10,000$, making the $H_{TCA}$ state essentially unpopulated. The fits are very poor. For the noncompetitive model, $K_C = 0.745 \mu M, K_T = 7.34 \mu M, L = 1.3$, and $y = 10,000$. B, normalized fluorescence anisotropy data for 8-fluo-cAMP binding to HCN2-CNBDxt fit with the partially competitive model at the top. Here, $x = 10^4$ resulted in a good fit to the data. For the competitive model, $K_C = 0.745 \mu M, K_T = 7.34 \mu M, L = 1.3$, and $x = 100$. The data are plotted as means ± S.E.
weakly dependent on TRIP8b concentration. However, the partially competitive model (green line) shows that the probability of finding the CNBD in the active state is strongly dependent on TRIP8b concentration. We were able to show that TRIP8b had little effect on the probability that HCN2-CNBDxt would undergo the conformational change into the active state, confirming again that a partial competition model best fits the data.

Figure 7. DEER data support the partial competition hypothesis. A, model predictions for the fraction of active CNBD molecules with 25 μM HCN2-CNBDx and 1 mM cAMP as a function of TRIP8b concentration for both the non-competitive and competitive models. For both models, $K_c = 0.745 \, \mu M$, $K_r = 7.34 \, \mu M$, and $L = 1.3$. For the noncompetitive model $y = 100$, and for the competitive model $x = 100$. The noncompetitive model (blue line) shows that the probability of finding the CNBD in the active state is strongly dependent on TRIP8b concentration. However, the partially competitive model (red line) shows that the probability of finding the CNBD in the active state is only weakly dependent on TRIP8b concentration. The vertical dotted line marks 300 μM TRIP8bcore. B, distance distributions of HCN2-CNBDxt SS63C/A624C (blue), with 300 μM TRIP8bcore (cyan), with 1 mM 8-fluoro-cAMP (red), and with 1 mM 8-fluoro-cAMP and 300 μM TRIP8bcore (green).

Adding a large amount of TRIP8bcore (300 μM) to HCN2-CNBDxt (25 μM) would have almost no effect on the conformational equilibrium between the resting and active states. On the other hand, if TRIP8b regulates HCN channels by a non-competitive mechanism, the model predicts that under that same condition, there would be a dramatic reduction in the probability that HCN2-CNBDxt would undergo the conformational change into the active state. DEER provided us with a powerful technique to look at conformational equilibria in purified proteins. By measuring DEER distributions under the conditions that we modeled, we were able to show that TRIP8b had little effect on the probability that HCN2-CNBDxt would undergo the conformational change into the active state, confirming again that a partial competition model best fits the data.

NMR work from two groups also shed light on the mechanism of TRIP8b inhibition (10, 11). Both groups used chemical shift perturbation analysis to look at residues on the CNBD of HCN channels. Although the two studies used different HCN isoforms (HCN2 and HCN4), the similarity in structures and inhibition by TRIP8b suggests that the interactions and mechanisms of inhibition will be similar (10, 11, 19). Overall, the two studies show similar chemical shift perturbations when TRIP8b is added to the CNBD of HCN. The residues that were most affected cluster in the C-helix, the cAMP-binding site, and the C-linker region, both regions that are involved in ligand binding and the subsequent conformational change. Interestingly, a new structure of the full human HCN1 channel shows two helices after adding a large amount of TRIP8bcore (300 μM) to HCN2-CNBDxt (25 μM) would have almost no effect on the conformational equilibrium between the resting and active states. On the other hand, if TRIP8b regulates HCN channels by a non-competitive mechanism, the model predicts that under that same condition, there would be a dramatic reduction in the probability that HCN2-CNBDxt would undergo the conformational change into the active state. DEER provided us with a powerful technique to look at conformational equilibria in purified proteins. By measuring DEER distributions under the conditions that we modeled, we were able to show that TRIP8b had little effect on the probability that HCN2-CNBDxt would undergo the conformational change into the active state, confirming again that a partial competition model best fits the data.
the cAMP-binding site or distorts the binding site by making interactions with the binding pocket, thus inhibiting the cyclic nucleotide dependence of these channel by acting like a partially competitive antagonist.

**Experimental procedures**

**Molecular biology**

All constructs used to express the HCN2 CNBD and TRIP8b fragments were subcloned into the pETM11 vector. The proteins were separated from the histidine tag by a tobacco etch virus (TEV) cleavable linker. TRIP8b<sub>core</sub> was derived from the murine TRIP8b(1a-4) isoform and contained residues 223–303 of that protein (9, 10). Murine HCN2-CNBDxt contained residues 488–640 (10). For EPR experiments, this construct was made in a cysteine-free background (10). 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 cDNAs encoding the full-length murine HCN2 channel in the pGHE vectors were kindly provided by Steven Siegelbaum and Bina Santoro (Columbia University) (21). The TRIP8b cDNAs were provided by Dane Chetkovich (Northwestern University) (3).

**Patch clamp recording**

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 (22). The vitelline membranes were manually removed, and currents were recorded in the inside-out patch-clamp configuration (23) with an EPC-10 patch-clamp amplifier (HEKA Elektronik). HCN2 channels are known to run down upon patch excision because of phosphatidylinositol 4,5-bisphosphate depletion (24). Thus, we allowed 20–30 min for the HCN current from the patch to stabilize before starting the experiment. Patch pipettes were pulled from borosilicate glass and had resistances of 0.4–0.6 MΩ after fire polishing. The pipette and bath solutions for HCN2 recordings were as follows: 130 mM KCl, 3 mM HEPES, and 0.2 mM EDTA (pH 7.2). TRIP8b<sub>core</sub> and cAMP were added to these solutions at the concentrations indicated. The solutions were perfused onto the patches using a flow low volume perfusion system (pH 7.2). TRIP8b<sub>core</sub> and cAMP were added to these solutions at the concentrations indicated. The solutions were perfused onto the patches using a μflow low volume perfusion system (ALA Scientific Instruments). Patches were held at 0 mV, and HCN2 currents were elicited by applying a series of 2-s voltage steps ranging from −70 to −140 mV followed by a 1-s voltage pulse to −40 mV. The data were analyzed using Igor (WaveMetrics) and MATLAB (MathWorks).

**Protein expression, purification, and spin labeling**

For each protein expressed, the construct was transfected into BL21(DE3) cells. 2-Liter cultures of cells were grown at 37 °C to an optical density of 0.6–0.8. The cells were then induced with 1 mM isopropyl-β-D-thiogalactopyranoside and grown overnight at 18 °C. After growth and expression, the cells were pelleted by centrifugation at 4,000 × g at 4 °C for 10 min and resuspended in 150 mM KCl and 30 mM HEPES at pH 7.4 (for HCN2-CNBDxt) or pH 8.5 (for TRIP8b<sub>core</sub>). DNase at a final concentration of 5 μg/ml and two tablets of protease inhibitors (Complete 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 × g at 4 °C for 45 min. The lysate was then purified on a Ni<sup>2+</sup> affinity resin column (HisTrap HP; GE Healthcare). For most experiments, the octahistidine tag was removed by TEV protease cleavage overnight at 4 °C. For the biolayer interferometry experiments, the octahistidine tag was left on the amino terminus of HCN2-CNBDxt. For anisotropy and DEER experiments, the proteins were then labeled with 100 μM monobromobimane (mBBr) or MTSL (Toronto Research Chemicals), respectively, 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 purified on an ion-exchange column. For HCN2 proteins, a cation-exchange column was used (HiTrap SP FF; GE Healthcare). For TRIP8b(1a-4) and for TRIP8b<sub>core</sub>, an anion-exchange column (HiTrap Q HP; GE Healthcare) was used. The proteins were eluted with a KCl gradient from 15 mM to 1 M. Fractions with protein were pooled and concentrated using a 3-kDa molecular mass cutoff centrifugal filter (Vivaspin; General Electric). The samples were then buffer-exchanged into a 150 mM KCl, 30 mM Tris, pH 8.4, solution using a PD-10 column (GE Healthcare). For CW EPR and DEER experiments, the protein was buffer-exchanged into D<sub>2</sub>O with 150 mM KCl, 30 mM Tris, pH 8.4, and 10% glycerol.

**Fluorescence anisotropy**

Fluorescence anisotropy was recorded using a Fluorolog 3 spectrofluorometer (Horiba, Jobin Yvon). The cAMP binding experiments utilized a fluorescent analog called 8-(2-[fluoresceinyl] aminoethylythio)adenosine-3′,5′-cyclic monophosphate (8-fluo-cAMP) (Biolog, Bremen, Germany). Anisotropy experiments with 390-nm (mBBr) or 494-nm (8-fluo-cAMP) (Biolog, Bremen, Germany). Anisotropy experiments were performed as previously described (25). To estimate binding affinity, plots of the anisotropy versus total HCN2-CNBDxt concentration were fit using the following first-order reaction scheme,

\[
R + A \leftrightarrow RA
\]

(\text{Eq. 1})

Anisotropy

\[
= \alpha \left[ \frac{(R_i + K_d + A_i)}{\sqrt{\left(R_i - K_d - A_i\right)^2 - 4AR_i}} \right] + \beta
\]

(\text{Eq. 2})

where \(R\), \(A\), and \(RA\) are concentrations of the free receptor, ligand, and receptor-ligand complex, respectively. \(R\) and \(A\) are total receptor and ligand concentrations, \(K_d\) is the dissociation constant, and \(\alpha\) and \(\beta\) are the scaling factor and offset factor, respectively.

**Biolayer interferometry**

Binding kinetics of TRIP8b<sub>core</sub> to the HCN2-CNBDxt were determined using the Octet Red 96 (ForteBio, Pall Life Sciences). Optical probes coated with Ni-NTA were loaded with an octahistidine-tagged HCN2-CNBDxt. All reaction solutions (200 μl) were loaded into black well-plates. The reaction
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buffer was 150 mM KCl and 30 mM HEPES, pH 8.4. There was no binding of TRIP8bcore to the unloaded probes. Binding kinetics of multiple concentrations of TRIP8bcore was measured simultaneously using eight probes. The data were acquired using the software for the Octet Red 96 and then analyzed offline using a script written for Igor (Wavemetrics). Binding kinetics for each concentration of TRIP8bcore were fit with the following equation:

\[ F = \alpha \times \left( e^{-(k_{on} + k_{off} + t)} \right) + \beta \]  
  (Eq. 3)

where \( F \) is the signal from the Octet Red 96, \( k_{on} \) and \( k_{off} \) are the binding and unbinding rate constants, respectively, and \( \alpha \) and \( \beta \) are the scaling factor and offset factor, respectively. The rate constants were fit globally using the global fit routine in Igor to come up with a single set of values that best fit the data at all the concentrations of TRIP8bcore. In addition, the binding kinetics of different concentrations of TRIP8bcore were fit with the following equation:

\[ k_{obs} = k_{on} \times [\text{TRIP8bcore}] + k_{off} \]  
  (Eq. 5)

as a second method for estimating \( k_{on} \) and \( k_{off} \).

EPR data collection and analysis

Doubly spin-labeled HCN protein for intraprotein DEER measurements was diluted to \( \sim 50 \mu M \). TRIP8bcore or cAMP was added as indicated in the text. 50 \( \mu L \) of each protein sample was inserted into a 1.65-mm outer-diameter quartz tube (Sutter, Q165-115-10). In an attempt to trap the room temperature equilibrium conformational ensemble, samples were flash frozen by rapid placement into liquid nitrogen. This procedure is known to result in a narrower ensemble of spin label rotamers (16, 17).

DEER data were acquired on a 33–35 GHz Bruker ELEXSYS E580 spectrometer with an overcoupled dielectric resonator (Bruker EN5107D2, 34.1 GHz, Q-factor 300–700). Experiments were performed at 60 K using a liquid-helium cooling system (Oxford). The four-pulse, dead-time free DEER sequence \( \left( (\pi/2)_{\text{probe}} - \tau_1 - (\pi)_{\text{probe}} - \tau_1 + t - (\pi)_{\text{pump}} - (\tau_2 - t) - (\pi)_{\text{pump}} - \tau_2 \right) \) 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 \( t \) 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 h.

DEER distance distributions were obtained using Deer-Analysis2013 (26). 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 bands shown in the distance distributions correspond to two standard deviations of the time domain noise. Molecular graphics and analyses were performed with UCSF Chimera and PyMOL (27–29).

Modeling of TRIP8bcore inhibition of cAMP binding to HCN2-CNBDxt

To distinguish between two mechanistic hypotheses for TRIP8b regulation of HCN channels, we created a six-state model to fit our binding data. The model is shown in Fig. 4 (B and C). The six states represent the following: \( H_U \) is the completely unbound HCN-CNBD, \( H_C \) is the cAMP bound HCN-CNBD in the resting conformation, \( H_{CA} \) is the cAMP bound HCN-CNBD in the active conformation, \( H_T \) is the TRIP8b bound HCN-CNBD, \( H_{TC} \) is the cAMP and TRIP8b bound HCN-CNBD in the resting state, and \( H_{TCA} \) is the cAMP and TRIP8b bound HCN-CNBD in the active state. A combination of fluorescence anisotropy, biolayer interferometry, and DEER experiments were used to determine binding affinities for cAMP and TRIP8b to HCN2-CNBDxt and to determine the equilibrium constant for the conformational change, \( L \). We then varied the parameter \( x \) or \( y \) (see Fig. 4, B and C) to produce the best fit of each model to the cAMP binding data with different concentrations of TRIP8bcore. Igor was used to calculate the roots of the following equations:

\[ H_{tot} = H_U + H_C + H_{CA} + H_T + H_{TC} + H_{TCA} \]  
  (Eq. 6)

\[ C_{tot} = C + H_C + H_{CA} + H_{TC} + H_{TCA} \]  
  (Eq. 7)

\[ T_{tot} = T + H_T + H_{TC} + H_{TCA} \]  
  (Eq. 8)

\[ K_C = \frac{H_U \times C}{H_C} \]  
  (Eq. 9)

\[ K_T = \frac{H_T \times T}{H_T} \]  
  (Eq. 10)

\[ L = \frac{H_{CA} \times H_{TC}}{H_C \times H_{TCA}} \]  
  (Eq. 11)

For the noncompetitive model,

\[ y = \frac{H_{CA} \times H_{TC}}{H_C \times H_{TCA}} \]  
  (Eq. 12)

For the competitive model,

\[ x = \frac{H_C \times H_T}{H_U \times H_T} \]  
  (Eq. 13)

where \( H_{tot}, C_{tot}, \) and \( T_{tot} \) are the total concentrations of HCN2-CNBDxt, cAMP, and TRIP8bcore, respectively, used in the experiment, and \( C \) and \( T \) are the concentrations of free cAMP, and TRIP8bcore, respectively. The other variables are defined in
Fraction cAMP bound = 1 - $\frac{C}{C_{tot}}$  \hspace{1cm} (Eq. 14)

**Statistics**

The data were plotted as means ± S.E. Student’s t test was used to determine significance at $p < 0.05$.

**Author contributions**—J. R. B. designed, conducted, analyzed experiments, and wrote the paper. H. A. D. designed, conducted, and analyzed experiments. W. N. Z. and S. S. analyzed experiments and wrote the paper.

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Mechanism for the inhibition of the cAMP dependence of HCN ion channels by the auxiliary subunit TRIP8b

John R. Bankston, Hannah A. DeBerg, Stefan Stoll and William N. Zagotta

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