An \( \alpha \)-\( \pi \) transition in S6 shapes the conformational cycle of the bacterial sodium channel NavAb

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Voltage-gated sodium channels play an important role in electrical signaling in excitable cells. In response to changes in membrane potential, they cycle between nonconducting and conducting conformations. With recent advances in structural biology, structures of sodium channels have been captured in several distinct conformations, which are thought to represent different functional states. However, it has been difficult to capture the intrinsically transient open state. We recently showed that a proposed open state of the bacterial sodium channel NavMs was not conductive and that a conformational change involving a transition to a \( \pi \)-helix in the pore-lining S6 helix converted this structure into a conducting state. However, the relevance of this structural feature in other sodium channels, and its implications for the broader gating cycle, remained unclear. Here, we propose a comparable open state of another class of bacterial channel from *Aliarcobacter butzleri* (NavAb) with characteristic pore hydration, ion permeation, and drug binding properties. Furthermore, we show that a \( \pi \)-helix transition can lead to pore opening and that such a conformational change blocks fenestrations in the inner helix bundle. We also discover that a region in the C-terminal domain can undergo a disordering transition proposed to be important for pore opening. These results support a role for a \( \pi \)-helix transition in the opening of NavAb, enabling new proposals for the structural annotation and drug modulation mechanisms in this important sodium channel model.

Introduction

Voltage-gated sodium ion channels (Navs) are membrane proteins that selectively transport sodium ions across cell membranes. They open in response to membrane depolarization, and, in excitable cells, are responsible for the fast-rising phase of the action potential. They are further characterized by rapid inactivation, during which the pore becomes nonconductive despite continued depolarized conditions (Ahern et al., 2016). Eukaryotic Nav channels are pseudotetrameric single-chain proteins consisting of around 2,000 residues. Bacterial sodium channels, on the other hand, have a simpler homotetrameric architecture, where each of four identical subunits comprises around 270 residues. As bacterial channels share important aspects of structure and function with eukaryotic Nav channels, they have been used as prototypes to understand structure–function relationships in this superfamily of channels (Payandeh and Minor, 2015).

Several structures of bacterial (Irie et al., 2018; Payandeh et al., 2011; Gamal El-Din et al., 2018; Jiang et al., 2018; Wisedchaisri et al., 2019; McCusker et al., 2012; Payandeh et al., 2012; Lenaeus et al., 2017; Tsai et al., 2013; Zhang et al., 2012; Bagnéris et al., 2014; Bagnéris et al., 2013; Sula et al., 2021; Naylor et al., 2016; Saït et al., 2020; Sula et al., 2017; Shaya et al., 2014; Arrigoni et al., 2016) and eukaryotic Nav channels (Jiang et al., 2020; Jiang et al., 2021a; Shen et al., 2018; Li et al., 2021; Pan et al., 2018; Shen et al., 2019; Jiang et al., 2021b) have been determined. A prototypical Nav channel consists of three types of domains—voltage-sensing domains (VSDs), a pore domain, and an intracellular C-terminal domain (CTD; Fig. 1A). Each VSD constitutes a four-helix bundle made of S1–S4, where S4 contains several arginine residues responsible for sensing the changes in potential difference across the membrane. Each channel contains four VSDs, each contributed by one subunit, that are arranged around a central pore. In their resting state, the VSDs’ S4 helices are stabilized in an “inward” or “down” position. Upon activation, the S4 helices transition into an “outward” or “up” position (Wisedchaisri et al., 2019). The pore domain consists of a tetrameric inverse cone-shaped arrangement of the S5 and S6 helices from each of the four subunits. The pore is thought to open and close at the so-called intracellular bundle crossing, where an expanded pore likely represents an open state and a constricted pore likely corresponds to a closed state. In the resting state of the channel, the VSDs are in a resting conformation and the pore is nonconductive. In the open state, the VSDs are activated and the pore is dilated. In the inactivated...
state, the VSDs remain activated, but the pore is nonconductive. In addition to these transmembrane domains, bacterial Nav channels possess a CTD that extends into the cytoplasm. This domain consists of two main regions—a neck region (proximal to the transmembrane region of the protein) and a distal coiled-coil region (Fig. 1 A). Disordering of the neck region of the CTD favors activation and pore opening in bacterial sodium channels, such as NavAe and NavSp (Shaya et al., 2014; Arrigoni et al., 2016). The CTD is also involved in channel inactivation through an unknown mechanism (Gamal El-Din et al., 2019).

Structures of the sodium channel from *Aliarcobacter butzleri* (NavAb) have been described as representing three functional states (Wisedchaisri et al., 2019; Payandeh et al., 2012; Lenaes et al., 2017). Based on these structures, the following functional cycle has been proposed (Fig. 1, B and C). Starting from a resting state (PDB ID 6P6W) with VSDs down and the pore in a constricted conformation, the channel transitions to a metastable preopen state in which the VSDs are activated and the pore is nonconductive. This transition may occur in two steps, first to a state with a highly constricted and symmetric pore (PDB ID 5VB2), followed by the second with a slightly wider and symmetric pore (PDB ID 3RVY). After this, the pore transitions to an open state that features an expanded pore (PDB ID 5VB8). Finally, the channel moves to an inactivated state wherein the VSDs are still activated but the pore is constricted and asymmetrical (C2 symmetry; PDB ID 4EKW). In 5VB2, an ordered
coiled-coiled CTD was resolved, but this domain is missing from all other structures.

Although this proposed cycle is consistent with some functional predictions, most structures of NavAb are made of constructs containing one or more mutations, and it is thus unclear how well they represent functional states of wild-type (WT) NavAb. The open state has been especially difficult to capture given the propensity of these channels to rapidly inactivate. Indeed, the structure of NavAb that is most likely to represent an open state, determined by x-ray crystallography, contains the pore-domain mutation, I217C, and is truncated after the S6 helix, thus deleting the CTD (Lenaeus et al., 2017). As molecular dynamics (MD) simulations indicated that the pore of this mutant was hydrated and conducted sodium ions, this structure was assigned as an open state (Lenaeus et al., 2017). However, the functional properties of this structure upon reversion to the WT sequence remained unclear.

In our recent study of the related bacterial channel NavMs (Choudhury et al., 2022), we found that a structure initially thought to represent an open state with an expanded pore and composed entirely of α-helical S6 helices was dehydrated and nonconductive to sodium ions. Introducing a short π-helical stretch in S6 in this channel resulted in the rotation of the S6 C-terminus and pore hydration, allowing sodium ion permeation in NavMs. In TRP channels, a π-helix formation and asparagine reorientation was recently proposed to be involved in pore opening (Kasimova et al., 2018; Kasimova et al., 2018b; Zubcevic and Lee, 2019). In addition, a reprocessing of the cryo-EM densities of Nav1.7 showed that the S6 helix in domain IV could be captured both in an α- and in a π-state (Huang et al., 2022). Inspired by these observations, we hypothesized that the opening of NavAb may also require a transition to the π-helix in S6.

Local anesthetic drugs act by binding to sodium channels (Fozzard et al., 2011), several of them in a state-dependent manner. Bacterial sodium channels are known to be modulated by these drugs in a similar manner as their eukaryotic counterparts, making them useful tools to study state-dependent drug binding (Payandeh and Minor, 2015; Lee et al., 2012). Depending on their protonation states, such drugs can access the Nav central cavity either via a hydrophobic pathway involving partitioning of the drugs in the membrane or via a hydrophilic pathway across the intracellular ion gate (Hille, 1977). Indeed, structures have revealed the presence of fenestrations that connect the Nav central cavity to the membrane (Payandeh et al., 2012). These fenestrations were confirmed to play an important role in a tonic block, whereby drugs access the central cavity when the pore is closed (Gamal El-Din et al., 2018). The generation of a new open model poses an opportunity to study the accessibility of local anesthetics to their binding sites in various states of NavAb.

In this study, we aimed to revise the proposed conformational cycle of NavAb and to describe detailed features of the pore domain in putative resting, open, and inactivated states of the channel. To do so, we first used MD simulations to describe the full-length structure of WT NavAb in a presumed open state. Reverting the I217C mutation dehydrated the pore, leading to a model that was impermeable to sodium ions. Given our previous model of NavMs pore opening (Choudhury et al., 2022), we then constructed an alternate model of the NavAb pore, in which introducing a π-helix in S6 results in a hydrated and conductive pore. This open-pore model was validated by sodium ion and local-anesthetic permeation free energy calculations. We then used the open model to characterize state-dependent properties of hydrophobic fenestrations and lidocaine binding. Finally, we posited a new opening mechanism for NavAb, in which the formation of a π-helix in S6 and disordering of the neck region of the CTD are prerequisites for pore expansion.

Materials and methods

Model building

Four different NavAb structures were used in this study (Fig. 1, B and C): a resting state (PDB ID 6P6W), in which the VSDs are at rest, the symmetric pore is contracted, and the CTD is truncated; a tentatively assigned open state (PDB ID 5VB8), in which the VSDs are activated, the symmetric pore is expanded, and the CTD is again truncated; and two decoupled states, tentatively assigned as preopen (PDB ID 5VB2) and inactivated (PDB ID 4EKW), in which the VSDs are activated and the pore is contracted. In 5VB2, the pore domain has C4 symmetry and the CTD is ordered, while in 4EKW the pore domain has only C2 symmetry (partly asymmetric) and the CTD is truncated. All the mutations in the structures were reverted to WT using CHARMM-GUI (Lee et al., 2019). Incomplete loops were also modeled using CHARMM-GUI (Lee et al., 2019).

Alternative models of NavAb with contracted or expanded pores were built by introducing a π-helix in the region of the S6 helix of 5VB2 and 5VB8, respectively. In all of these models, the VSDs remain activated. This was done using MODELLER 9.22 (Webb and Sali, 2016), following a protocol implemented in previous work on NavMs (Choudhury et al., 2022). Briefly, this consisted in shifting the structural alignment in S6 of the π-versus template α-models to introduce an unpaired backbone carbonyl. Specifically, we introduced one gap in the template (α) sequence prior to T1206, and a second in the target (π) sequence following D1219, thus shifting the region one position upstream from T1206 to the inner end of S6, resulting in the formation of a π-helix around T1206. This resulted in rotation of the pore helix following the π-helix around T1206. As a result of this, residues facing the pore in the α-model rotate away from it in the π-model, while a different set of residues—including N1211—now face the pore (Choudhury et al, 2022). Since all the residues in this region except N1211 are hydrophobic, the protonation states of the residues are not affected by the rotation.

For pore-opening adiabatic bias MD (ABMD; see below for details) simulations, we also built contracted-α and contracted-π NavAb models from PDB ID 5VB2 with the CTD deleted or by following the procedure described above. This approach ensured that the protein length was consistent with the expanded-pore models.

System preparation

Each model was embedded in a homogenous lipid bilayer consisting of 400 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine
(POPC) using the CHARMM-GUI Membrane builder (Lee et al., 2019). Lidocaine-protein complexes were prepared by randomly placing the drug in the central cavity of the different models. Each system was hydrated by adding ~45 Å layers of water to either side of the membrane. Lastly, the systems were neutralized with 150 mM NaCl. The Charmm36 force field was used to describe the interactions between proteins (Huang and MacKerell, 2013), lipids (Klauda et al., 2010), and ions. In addition, we considered additional simulations using Charmm36m to study CTD disordering, as this particular forcefield is better suited to describe disordered proteins (Huang et al., 2017). Nonbonded fixes (NBFix) were considered in the description of interactions of sodium ions with carboxyl and carbonyl groups (Luo and Roux, 2010). The TIP3P model was used to describe all water molecules (Jorgensen et al., 1983). Parameters for charged lidocaine were taken from our previous study (Choudhury et al., 2022).

**Simulation details**

For the expanded-pore states (with or without lidocaine) and the protein–lidocaine system simulations, equilibration was done following the protocol reported in Lenaeus et al. (2017). Each system was minimized for 5,000 steps using the steepest descent and equilibrated with a constant number of particles, pressure, and temperature for 30 ns, during which position restraints on the protein were gradually released. Initially, the heavy atoms of protein were restrained with a force constant of 1,000 kJ/mol/nm² for 10 ns. Then, the protein backbone was restrained with a force constant of 1,000 kJ/mol/nm² for 10 ns. Finally, the protein Ca atoms were restrained with a force constant of 1,000 kJ/mol/nm² for 10 ns. For all the contracted models, equilibration was carried out in six steps according to the default CHARMM-GUI protocol, extending the duration of each step (the first three steps were run for 2 ns each and the last three steps for 10 ns each; Lee et al., 2019). During both equilibration and production, a time step of 2 fs was used, the pressure was maintained at 1 bar through Berendsen pressure coupling, and the temperature was maintained at 300 K through Berendsen temperature coupling (Berendsen et al., 1984) with the protein, membrane, and solvent coupled, and the LINCs algorithm (Hess et al., 1997) was used to constrain the bonds involving hydrogen atoms. For long-range interactions, periodic boundary conditions and particle mesh Ewald were used (Darden et al., 1993). For short-range interactions, a cut-off of 12 Å was used. Production simulations were run with or without restraints for different systems, as shown in Table 1 using Parinello-Rahman pressure coupling (Parinello and Rahman, 1981) and Nosé-Hoover temperature coupling (Hoover, 1985). Simulations were performed using GROMACS 2020.3 (Abraham et al., 2015; Lindahl et al., 2021). Some simulations were performed under a transmembrane voltage of ±750 mV via application of an external electric field. The transmembrane voltage is calculated as the product of the electric field and the simulation box length along which the field is applied. Such a transmembrane voltage did not cause any disruption to the protein/membrane over the relatively short simulation time considered.

The free energy profiles of sodium ions and lidocaine permeation along the pore axis were calculated using the accelerated weight histogram (AWH) method (Lindahl et al., 2014), as shown in Table 2. In brief, for each equilibrated structure, we applied one independent AWH bias and simulated for 200 ns each, with six walkers sharing bias data and contributing to the same target distribution. The target distribution chosen was a flat (uniform) distribution. Each bias acts on the z-axis defined using the center-of-mass z-distance between one central sodium ion/lidocaine (all atoms) and all atoms of S1178 residues in NavAb, with a sampling interval corresponding to the entire box length. The rate of change of each AWH bias was initialized by setting the initial average free energy error to 20 kJ/mol, and the diffusion constant was set to 0.00005 nm²/ps. To keep the solute close to the pore axis, the coordinate radial distance was restrained to stay below 10 Å by adding a flat-bottom umbrella potential. All the Ca atoms of the protein were restrained by imposing harmonic potentials with force constants of 1,000 kJ mol⁻¹ nm⁻² to remain as close as possible to the experimental structures. The average potential of mean force (PMF) profile and the error bars were calculated from a single walker since the walkers communicate with each other. For all systems, except the expanded-π-model with lidocaine, the average PMF profile was calculated by taking the data from the last 100 ns in intervals of 10 ns, and the error bars were calculated by calculating the standard deviation. For the expanded-π-model with lidocaine, data from the last 50 ns at 10-ns intervals were chosen for calculating the average and standard deviation.

Additionally, the lidocaine-permeation-free energy along the pore axis was also calculated using multiple walker well-tempered metadynamics (Laio and Parrinello, 2002; Barducci et al., 2008) in GROMACS 2020.5 patched with PLUMED 2.7.3 (Bonomi et al., 2009; Tribello et al., 2014), as shown in Table 2. Each bias acts on the z-axis defined using the center-of-mass z-distance between lidocaine (all atoms) and all atoms of S1178 residues in NavAb. The Gaussian’s height and width were set at 1.2 kJ/mol and 0.05 nm, respectively. Upper and lower restraints (as shown in the corresponding x axes of the free energy plots) were applied to restrict the sampling space. For the models with constricted pores, the sampling was restricted to the central cavity of the pore. To keep the drug close to the pore axis, the coordinate radial distance was restrained to stay below 10 Å by restraining the x and y component of the center-of-mass distance. The free energy profile was estimated using the sum_hills tool in PLUMED. The average free energy profile was calculated by taking the data from the last 30 ns in intervals of 5–10 ns, and the error bars were estimated by calculating the standard deviation.

**ABMD for pore opening**

The pore opening was studied by performing ABMD (Marchi and Ballone, 1999) in GROMACS 2020.5 patched with PLUMED 2.7.3 (Bonomi et al., 2009; Tribello et al., 2014) as shown in Table 2. ABMD is a simulation method in which whenever the system moves closer toward the target system along the collective variable, the harmonic potential is moved to this new position, resulting in enhancing the system’s sampling toward the target value of the collective variable. The bias potential was
applied along the distance between the center of masses of Ca atoms of I1216/I1217 and I1119 (for the α-model and the π-model, respectively). The force constant for the harmonic potential was set at 5,000 kJ/mol/nm², while the target value for the collective variable was set at 1 nm. The simulations were started from a contracted α- and π-model and were stopped when the system reached the expanded states. ABMD also outputs the square of force applied along the distance between the center of masses of C atoms of I1216/I1217 and I1119 (for the α-model and the π-model, respectively).

The expanded-α-structure of NavAb is nonconductive

To test the relevance to the WT channel function of the proposed open-state structure of NavAb, in which the VSDs were captured in the open-state structure of NavAb, in which the VSDs were captured.

Table 1. Summary table of equilibrium simulations performed

| System | Equilibration | Production |
|--------|---------------|------------|
| NavAb expanded α-model (PDB ID 5VB8) | 30 ns | 185 ns (unrestrained) |
| NavAb expanded α-model I217C (PDB ID 5VB8) | 30 ns | 250 ns (Ca restrained) |
| Expanded π-model | 30 ns | 154 ns (unrestrained) |
| NavAb expanded α-model (PDB ID 5VB8) | 30 ns | 200 ns (Ca restrained) |
| NavAb expanded α-model (+750 mV) | 30 ns | 200 ns (Ca restrained) |
| NavAb expanded α-model (−750 mV) | 30 ns | 200 ns (Ca restrained) |
| NavAb expanded π-model (PDB ID 5VB8) | 30 ns | 154 ns (unrestrained) |
| NavAb expanded α-model + lidocaine | 30 ns | 200 ns (Ca restrained) |
| NavAb asymmetric α model + lidocaine | 30 ns | 200 ns (unrestrained) |
| NavAb resting α model + lidocaine | 30 ns | 200 ns (unrestrained) |
| NavAb expanded π-model + lidocaine | 30 ns | 200 ns (Ca restrained) |
| NavAb expanded π-model N1211L | 30 ns | 200 ns (Ca restrained) |
| NavAb contracted α-model (PDB ID 5VB2) with CTD—300 K—Charmm36m | 36 ns | 200 ns (unrestrained) |
| NavAb contracted α-model (PDB ID 5VB2) with CTD—320 K—Charmm36m | 36 ns | 308 ns (unrestrained) |
| NavAb contracted α-model (PDB ID 5VB2) with CTD—360 K—Charmm36m | 36 ns | 309 ns (unrestrained) |
| NavAb contracted α-model (PDB ID 5VB2) with CTD—400 K—Charmm36m | 36 ns | 215 ns, 223 ns, 232 ns (unrestrained) |
| NavAb contracted π-model (PDB ID 5VB2) with CTD | 36 ns | 200 ns (unrestrained) |
| NavAb contracted π-model (PDB ID 5VB2 WT) without CTD | 36 ns | Equilibrated structure used for ABMD simulation × 3 |
| NavAb contracted α-model (PDB ID 5VB2) without CTD | 36 ns | Equilibrated structure used for ABMD simulation × 3 |
| NavAb contracted α-model (PDB ID 5VB2 N1211L) without CTD | 36 ns | Equilibrated structure used for ABMD simulation × 3 |

Restrained production simulations (where indicated) were used to prevent pore collapse in the absence of CTD. Ion permeation and drug permeation free energy calculations.

Analysis

The water number density, water hydration-free energy, and pore radius were calculated using the channel annotation package (CHAP; Klesse et al., 2019). For the water density analysis, frames were extracted every 100 ps from the trajectories of the respective models. For the fenestration radius analysis, 200 equally spaced frames were extracted from the trajectories of the expanded/contracted α- and π-models. We calculated the time-averaged radius for each fenestration in each model. Since all the models are homotetrameric, we calculated the average of the four-fenestration radius and the uncertainty on the average estimate, as the average of the standard deviation, by taking the square root of the average of the squares of the standard deviations. Representative binding poses of lidocaine were obtained by performing clustering on frames extracted from the AWH calculations corresponding to free energy minima. Clustering was done using gmx cluster, with a cutoff of 0.1 nm. The center of the most populated cluster was then chosen as the representative binding pose. A similar approach was used to extract the most representative structures from the pore-opening free energy profiles. Hydrogen bonding between N1211 and F1207 was calculated using gmx hbond. The RMSD for N1211 was calculated by first aligning the structures on their backbone atoms using gmx rms. The files necessary to reproduce the data herein can be found at https://osf.io/u8fg9/.

Online supplemental material

There are 18 supplemental figures that accompany this paper.

Results

The expanded-α-structure of NavAb is nonconductive

To test the relevance to the WT channel function of the proposed open-state structure of NavAb, in which the VSDs were captured...
in an activated state and the pore in an expanded conformation (PDB ID 5VB8), we first reverted the I1217C mutations and carried out equilibrium MD simulations, all in absence of the CTD.

The introduction of this π-helix resulted in the rotation of a conserved asparagine, N1211, toward the NavAb pore lumen (Fig. 2 B). Mutating this pore-facing asparagine to a hydrophobic residue (N1211L) did not affect pore hydration (Figs. S2 and S5), but slightly increased the barrier to sodium-ion permeation at the site of mutation (Fig. 2 D). N1211 also likely plays an important role in stabilizing the π-helix by forming a hydrogen bond with the backbone carbonyl of F1207. Indeed, N1211 is less mobile in the expanded-π-model than in the expanded-α-model (Fig. S6). Taken together, these results are consistent with the expanded-π-model representing a functional open state, and with N1211 possibly contributing to sodium-ion permeation.

Local anesthetic binding poses and access pathways vary with functional state

We hypothesized that a structural transition in S6 may impact the accessibility of neutral drugs to their binding site in the pore cavity through the pore-lining fenestrations. To test this, we calculated the time-averaged radius of the four fenestrations in the contracted-α, contracted-π, expanded-α, and expanded-π-models of NavAb. Comparing the fenestration radii for these models, we found that the π-models appear to have narrower fenestrations than the α-models (Fig. 3 A and B). Indeed, the fenestrations are obstructed by large hydrophobic residues F1207 and I1210 in both the contracted- and expanded-π-models (Fig. 3 B). This observation indicates that hydrophobic drugs can access the central cavity only in the context of a fully α-helical S6, irrespective of pore diameter.
We next investigated state-dependent drug access to the central cavity via the hydrophilic pathway by calculating lidocaine permeation-free energies across the channel gate in different functional states of NavAb. We particularly compared our presumed open expanded-π-model to two previously reported structures, one with resting VSDs and a symmetrically contracted pore (assigned resting; PDB ID 6P6W), and the other one with activated VSDs and an asymmetric conformation of the pore (assigned inactivated; PDB ID 4EKW). Unlike the expanded-π-model, the resting and inactivated models featured high barriers to drug access from the intracellular side (Fig. 3 C; and Figs. S9, S10 B, S11, S13, S14, and S15). In all three models, free-energy profiles indicated that the drug binds in the central cavity close to the selectivity filter, forming contacts with T1206 in both the resting and expanded-π-models. However, in the resting model, lidocaine was oriented parallel to the pore axis with its charged group pointing toward the selectivity filter, while in the expanded-π-model it was oriented parallel to the membrane plane (Fig. 3 D). This difference could be due to its presence in the expanded-π-model of the hydrophilic sidechain of N1211, which accommodates the charged group of lidocaine. Note however that lidocaine binding does not affect the flexibility of N1211 (Fig. S6). In the asymmetric inactivated model, the drug was oriented perpendicular to the membrane plane and shifted further toward the intracellular end of the pore, closer to V1213 (Fig. 3 D). These results indicate that there is a change in the binding pose of lidocaine as the channel transitions between different functional states and that the drug can access its binding site via the intracellular gate exclusively in the π model.

Neck region of the C-terminal domain can undergo disordering

Experimental structures and electrophysiology recordings have indicated a disordered segment in the neck region of the CTD in certain states of several bacterial sodium channels (Arrigoni et al, 2016). This conformational change is proposed to be important for pore opening. To test whether this region can undergo disordering in NavAb, we subjected the full-length constricted-α-structure of NavAb (PDB ID 5VB2) to unrestrained simulations at elevated temperatures (300 K, 340 K, 360 K, and 400 K, the latter in triplicate). Simulations at 320 K and 360 K showed limited disordering in the CTD (Fig. 4 B and C), while those at 400 K showed a complete disordering of the neck region (Fig. 4 D) spanning residues Q1226 to I1243 (Fig. 4 D). The secondary structure was overall preserved in the rest of the protein.
at 400 K (Fig. 4 D), suggesting that the CTD neck is specifically prone to losing its secondary structure.

A π-helix transition results in pore opening

The results above suggested that a π-helix in S6, along with an expanded pore, results in an open state of NavAb. In contrast, a contracted and entirely α-helical S6, as resolved in the closed experimental structure, results in an obstructed pore. Pore opening, therefore, involves an α- to π-helix transition in S6, on top of pore expansion. To determine whether the α- to π-helix transition occurs before or after the expansion of the pore, we characterized the force needed to open the pore in presence of an α- and π-helix. In the contracted-α-model, N1211 is pointing away from the pore into the peripheral cavity, attracting water from the cytoplasm into peripheral cavities (Fig. 5 A). We propose that these water-filled peripheral cavities, as well as the attractive interaction between hydrophobic pore-facing residues at the activation gate, could impede pore opening. In the contracted-π-model, on the other hand, N1211 faces the pore lumen rather than the peripheral cavities, which are hence no longer hydrated (Fig. 5 B). These observations could constitute a rationale for the low propensity of the α-model to expand.

To add further quantitative support to this rationale, we performed ABMD simulations to open the pore starting from the contracted-π- and α-model, using the distance between Ca atoms of I1216 (α-model)/I1217 (π-model) and I1119 across the four subunits (Fig. 5 C) as the collective variable. The collective variable was chosen because these residues are located at the intracellular ends of S6 and the S4–S5 linker, respectively, such that the distance between them falls dramatically during the transition from the preopen/contracted to open states of the activated channel. This collective variable indeed enabled...
sampling of expanded-pore conformations from simulations initiated from contracted pores (Fig. 5, D and E; and Figs. S16, S17, and S18) and did not lead to substantial helix unwinding or other nonphysical transitions (Fig. 5, D and E, insets). We performed three replicates, each starting from the contracted-\(\pi\)- and \(\alpha\)-models, and calculated the average total force across the replicates. The average total force for pore opening was lower when the simulations were initiated from the contracted-\(\pi\)-model relative to the contracted-\(\alpha\)-model, indicating that the \(\pi\)-model is more likely to open than the \(\alpha\)-one (Fig. 5 F). To further characterize whether the presence of the conserved Asn contributes to lowering the energy needed to open the \(\pi\)-model, we repeated the same simulations in the presence of the N1211 mutation, thus removing the only hydrophilic residue in this region. The average total force for pore opening increased in N1211L relative to the WT \(\pi\)-model (Fig. 5 D), indicating that, indeed, N1211 contributes to a favorable opening of the pore, and possibly rationalizing its conservation across Nav channels.

**Discussion**

Our first observation in this work was that the expanded-\(\alpha\) experimental structure of NavAb does not convincingly model a functional open state. Indeed, in simulations of this model, the pore was dehydrated, preventing sodium ion permeation across the gate and lidocaine access to its binding site in the central cavity. However, introducing a short \(\pi\)-helix in S6 produced a consistently hydrated pore, which allowed sodium/lidocaine permeation. One contributor to this phenomenon is likely the conserved asparagine (N1211), where mutation to leucine slightly increased the free-energy barrier to ion permeation.

What, then, is the mechanism of transition between the resting/closed and the activated/open states? We found that the CTD of NavAb has a region proximal to the membrane that can undergo a disordering transition. As similar disordering is thought to facilitate pore opening in other bacterial channels, a similar mechanism may contribute to the opening in NavAb. Indeed, our understanding of the involvement of the CTD in the gating mechanism had been hampered by the lack of resolution in that...
domain in most structures of NavAb. Conducting simulations at high temperatures revealed a propensity for the neck region to disorder, in agreement with observations in other bacterial channels (Arrigoni et al., 2016). Although not the focus of this study, future experimental or computational work—for example, temperature-based replica exchange simulations—could give us insights into the free energy needed to unfold this region in the various pore states, contributing to our understanding of the role played by this region.

Our pore-opening studies further showed that it is unfavorable to expand a constricted pore with an α-helical S6. Thus, channel activation could include a transition from α- to π-helix in S6 via a state comparable to our contracted-π-model. Indeed, the eukaryotic channel NavPas was recently captured in a conformation consistent with such a preopen functional assignment, with activated VSDs, a contracted pore, π helices in all of its S6 segments, and a dissociated inactivation particle (Shen et al., 2018; Shen et al., 2017). The symmetric expanded-α-model of NavAb, which was initially assigned as open (Lenaeus et al., 2017), may instead represent an inactivated state of the WT channel, possibly contributing to a multistep inactivation process along with the asymmetric contracted-α-structure (Payandeh et al., 2012). Resolving this complex inactivation mechanism thus remains an area of active research since the order in which π to α transitions, pore contraction, and possible changes in symmetry occurs is currently unknown. It has been observed in TRPV1 channels that a conformation with a hydrated pore usually features dehydrated peripheral cavities between the S6 and S4–S5 helices, whereas conformations with a dehydrated pore have hydrated peripheral cavities (Kasimova et al., 2018). In the NavAb contracted-α-model, N1211 attracts water into the peripheral cavities, apparently impeding pore opening. In the contracted-π-model, on the other hand, N1211 faces the pore lumen, and stabilizing hydrophobic interactions at the gate are no longer present. This remodeling appears to favor the opening of the hydrophobic activation gate, as witnessed in the pore-opening ABMD simulation of the π-model and the N1211L mutant.

Putting together these results, we propose a modified mechanistic picture for the functional cycle of bacterial sodium channels (Fig. 6). Five important processes occur during the transition between the resting and open state—the VSDs activate, an α- to π-helix transition occurs in S6 resulting in rotation of the inner half of the helix, peripheral cavities dehydrate, the
CTD disorders, and the pore expands. Activation of the VSDs in response to membrane depolarization is presumed to be the initiating step in this cycle. Then, we propose that the $\alpha$ to $\pi$ transition occurs, resulting in dehydration of peripheral cavities, finally followed by pore expansion. We also propose that the CTD disorders during the opening transition, though our data do not definitively indicate at what stage this occurs. We hypothesize that the energy to overcome barriers associated with $\pi$-helix formation, CTD disordering, and pore opening is contributed at least in part by coupling to VSD activation, which is energetically favorable, thanks to membrane depolarization. Nevertheless, as the mechanism of VSD-pore coupling in Nav channels is still being investigated by many groups (Carvalho-de-Souza and Bezanilla, 2019; Lu et al., 2002; Cowgill and Chanda, 2021), it is currently challenging to devise a quantitative mechanism.

Finally, we considered what aspects of the proposed mechanism for bacterial-Nav gating and inactivation may be applicable to eukaryotic channels. Indeed, we have now reported the transition of a region of S6 to a $\pi$-helix as a factor in the opening of two bacterial channels, NavMs and NavAb. In addition, the opening of certain TRP channels appears to involve an $\alpha$-to-$\pi$-helix transition and a reorientation of a conserved Asn residue (Kasimova et al., 2018; McGoldrick et al., 2018; Singh et al., 2018a; Zubcevic et al., 2018a; Cao, 2020), raising the possibility that this is a conserved mechanism across evolution. Eukaryotic sodium channels also possess a conserved asparagine residue (N1211) in their S6 helices, and structures have revealed the presence of $\pi$-helices in certain states of some isoforms. Most human sodium channels feature a $\pi$-helix in domains I and II, while the other two S6 helices (domains III and IV) are $\alpha$-helical (Jiang et al., 2020; Pan et al., 2018; Shen et al., 2019; Jiang et al., 2021b). Recently, refined analysis of Nav1.7 densities showed a transient $\pi$-helix in domain IV (Huang et al., 2022), indicating that this transition is indeed energetically possible. This conformational change has been linked to inactivation, posing provocative possibilities for future study of helical transitions in the gating cycle of both bacterial and eukaryotic channels. Another interesting consequence of $\pi$-helix formation in S6 was observed at the so-called fenestrations, which are thought to provide pathways for hydrophobic drugs, such as lidocaine or benzocaine, to access binding sites in Nav-channel pores (Gamal El-Din et al., 2018).

Figure 6. Proposed functional cycle for NavAb. Starting from the resting state with $\alpha$-helical S6, NavAb transitions to a preopen state with VSDs activated and $\alpha$ to $\pi$ transition. This conformation allows for favorable pore opening. Pore opening also involves CTD disordering; although the point at which this happens remains unclear. Inactivation involves the transition from $\pi$ to $\alpha$, but whether the pore is expanded, contracted in a symmetric or asymmetric manner, or even transitions between different conformations, remains unclear. Return to the resting state involves the deactivation of the VSD.
More precisely, hydrophobic drugs have been proposed to freely access the pore in closed states (Gamal El-Din et al., 2018). In NavAb, we found that the fenestrations are closed in π models irrespective of the dimensions (constricted/expanded) of the pore, while they are open in α-models. This finding is in agreement with a functional assignment of the eukaryotic channels NavPas and NALCN as preopen (Shen et al., 2018; Shen et al., 2017; Kschonsak et al., 2020), in which all S6 segments contain π helices, and the fenestrations are obstructed. Moreover, recent structures of human Nav1.7 indicated that the presence of a π-helix in subunit IV closed the fenestration at the interface with subunit I (Huang et al., 2022). Lidocaine binding to different functional states of NavAb revealed that the drug binds near the selectivity filter in resting and π-models, in agreement with previous simulations in NavMs and NavPas, respectively (Gamal El-Din et al., 2018; Buyan et al., 2018). Lidocaine binds nearer the intracellular gate in the expanded-π model, in agreement with multi-level parallelism from laptops to supercomputers.

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K. Choudhury performed the simulations and analyzed the results. K. Choudhury, R.J. Howard, and L. Delemotte interpreted the data and wrote the manuscript.

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Pore opening mechanism in NavAb

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Figure S1. **Pore water density along time for different models.** *(A)* Water number density profile over time along the central pore axis of WT expanded-α-model of NavAb at 0 mV. *(B)* Water number density profile over time along the central pore axis at +750 mV. *(C)* Water number density profile over time along the central pore axis at −750 mV. The activation gate is transiently hydrated. *(D)* Water number density profile over time along the central pore axis in the I1217C mutant at 0 mV.
Figure S2. Time-averaged water number density plot along the central pore axis for various NavAb models. Standard deviation shown in light shades of respective colors for different models.

Figure S3. Convergence of free energy profile of sodium ion permeation in the NavAb expanded-α-model as determined using AWH. (A) Free energy profile at different times. The free energy was calculated across six walkers sharing the bias. (B) Target distribution at different times.
Figure S4. Convergence of free energy profile of sodium ion permeation in NavAb expanded-π-model (WT) as determined using AWH. (A) Free energy profiles at different times. (B) Target distribution at different times.

Figure S5. Convergence of free energy profile of sodium ion permeation in NavAb expanded-π-model (N1211L) as determined using AWH. (A) Free energy profiles at different times. (B) Target distribution at different times.
Figure 56. **Interactions and fluctuations of the conserved Asparagine (N1211) in the expanded-π-model.** (A) Backbone carbonyl of F1207 forming a hydrogen bond with the side chain of N1211, hence stabilizing the π-helix. (B) Number of hydrogen bonds between the F1207 backbone carbonyl and side chain of N1211 in the π-helix model as a function of simulation time. (C) RMSD of N1211 in different models. Expanded-α-model is in gray, expanded-π-model is in red, and expanded-π-model with lidocaine is in green.
Figure S7. Lidocaine permeation free energy profiles as determined using AWH for the expanded-α-model (black) and expanded-π-model WT (red) along the central pore axis of the lower half of the S6 helix. Cartoon representations in the background show the lower half of the S6 helix.
Figure S8. Unrestrained simulations of expanded pores result in pore collapse. (A and B) Structural alignment between the final model of the restrained simulations (A, red) and the unrestrained simulations (pink) shows that, in the absence of restraints, the pore collapses within 154 ns to a more contracted radius (pink), leading to dehydration (blue surface) around the constriction point (B).
Figure S9. Lidocaine permeation free energy profiles as determined using AWH for the resting model (black), expanded-\(\pi\)-model (red) and asymmetric \(\alpha\)-model (blue) along the central pore axis of the lower half of the S6 helix.

Figure S10. Convergence of free energy profile obtained using multiple walker well-tempered metadynamics for lidocaine permeation in NavAb expanded-\(\alpha\)-model. (A) Free energy profiles at different times. (B) Convergence of free energy profile of lidocaine permeation in NavAb expanded-\(\pi\)-model.
Figure S11. Convergence of free energy profile obtained using multiple walker well-tempered metadynamics for lidocaine permeation in NavAb expanded-α-model. (A) Free energy profiles at different times. (B) Convergence of free energy profile of lidocaine permeation in NavAb asymmetric α-model.

Figure S12. Convergence of free energy profile of lidocaine permeation in NavAb expanded-α-model as determined using AWH. (A) Free energy profiles at different times. (B) Target distribution at different times.
Figure S13. Convergence of free energy profile of lidocaine permeation in NavAb expanded-π-model as determined using AWH. (A) Free energy profiles at different times. (B) Target distribution at different times.

Figure S14. Convergence of free energy profile of lidocaine permeation in NavAb resting model as determined using AWH. (A) Free energy profiles at different times. (B) Target distribution at different times.
Figure S15. Convergence of free energy profile of lidocaine permeation in NavAb asymmetric α-model as determined using AWH. (A) Free energy profiles at different times. (B) Target distribution at different times.

Figure S16. Time evolution of the pore opening ABMD simulation starting from the contracted π model (WT) for two (A and B) of the replicas (the other replica is found in Fig. 5).
Figure S17.  
Time evolution of the pore opening ABMD simulation starting from the contracted α-model for two (A and B) of the replicas (the other replica is found in Fig. 5).

Figure S18.  
Time evolution of the pore opening ABMD simulation starting from the contracted π model (N1211L) for three replicas (A, B, and C).