

Local anesthetic and antiepileptic drug access and binding to a bacterial voltage-gated sodium channel

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Voltage-gated sodium (Na_v) channels are important targets in the treatment of a range of pathologies. Bacterial channels, for which crystal structures have been solved, exhibit modulation by local anesthetic and anti-epileptic agents, allowing molecular-level investigations into sodium channel-drug interactions. These structures reveal no basis for the “hinged lid”-based fast inactivation, seen in eukaryotic Na_v channels. Thus, they enable examination of potential mechanisms of use- or state-dependent drug action based on activation gating, or slower pore-based inactivation processes. Multimicrosecond simulations of Na_vAb reveal high-affinity binding of benzocaine to F203 that is a surrogate for FS6, conserved in helix S6 of Domain IV of mammalian sodium channels, as well as low-affinity sites suggested to stabilize different states of the channel. Phenytoin exhibits a different binding distribution owing to preferential interactions at the membrane and water-protein interfaces. Two drug-access pathways into the pore are observed: via lateral fenestrations connecting to the membrane lipid phase, as well as via an aqueous pathway through the intracellular activation gate, despite being closed. These observations provide insight into drug modulation that will guide further developments of Na_v inhibitors.

bacterial sodium channel | drug binding

Voltage-gated sodium (Na_v) channel inhibitors can modulate sensory or motor activity without threatening vital bodily functions, enabling a wide range of therapeutic applications. In particular, the ability to control Na_v channel current allows for effective use as local anesthetic (LA) and antiepileptic (AE) drugs, normalizing function in conditions of hyperexcitability, such as epilepsy, cardiac arrhythmias, hyperalgesia, and myotonia (1, 2). Recent breakthroughs in the solution of atomic structures of the bacterial Na_vBac family (3–6) provide an excellent opportunity to describe drug binding at the molecular level.

Pharmacological Na_v channel modulation is a complex phenomenon involving voltage- and state-dependent block of current, as well as prolongation of recovery from inactivated states, with interrelated observations that have implicated different active sites, pathways and inactivated channel states (7). Despite historical focus on fast inactivation (8), it has now been discovered that slow inactivation is affected by LA/AE drugs (9) and that mutation of key residues involved in inhibition impairs slow inactivation (10). The Na_vBac family lacks the Domain (D) III–DIV linker responsible for fast inactivation (3), enabling molecular-level investigation into this common modulation process. Moreover, key binding regions are well conserved in bacterial and mammalian channels (3, 11), and it has been shown that bacterial channels interact with common inhibitory drugs (11–14). However, questions remain as to how structurally disparate inhibitor molecules can cause qualitatively similar modulation of different mammalian and bacterial Na_v channels. Here, we explore the binding of LA benzocaine (BZC) and AE phenytoin (PHT) to the bacterial Na_vAb channel (3) (Fig. 1), providing detailed insight into Na_v modulation mechanisms.

Functional studies have suggested at least two major LA/AE binding sites: a high-affinity, state- and voltage-dependent site; and a low-affinity, less state-dependent site (15). The high-affinity site

has been the focus of considerable investigation (16) and appears to be shared by many drugs (2). Studies of mammalian Na_v have suggested this site is on helix S6 of DIV, involving residue FS6 [F1764 in Nav1.2 (16), F1579 in Nav1.4, F1759 in Nav1.5 (17)], as well as a Tyr (Y1771 in Nav1.2, Y1586 in Nav1.4, Y1766 in Nav1.5). Binding at this site is use-dependent, such that under brief repetitive stimuli that open the channel, the channel enters a stable nonconducting state, with a cumulative reduction of current (18), signifying preferential binding to an inactivated state (10). Binding to FS6 appears to involve cation- π interaction for charged lidocaine (19), but π -stacking for neutral agents such as BZC and PHT. In contrast, little information exists to pinpoint the low-affinity site(s), whose binding may result from a “combination of hydrophobic interactions” occurring when the channel is maintained at hyperpolarized voltages (15). Complete sampling of the drug-channel binding distribution is needed to define these binding modes.

Residue FS6 on domain IV of the mammalian channels is a critical feature in drug modulation, but does not exist at that position in the bacterial channels (Fig. 2A; with full comparison in *SI Appendix*, Fig. S1). The absence of this important residue may contribute to low LA affinities observed for NaChBac (12). However, the recently solved Na_vAb (3) possesses a nonconserved F203, which appears to mimic FS6. Fig. 2B shows a simple overlay of the hNa_v1.2 sequence on the Na_vAb structural template, with FS6 location and orientation similar to F203 (shifted only slightly along the S6 helix), and thus could play a similar role in drug binding. Furthermore, mammalian Na_v channels possess a second Phe in domain I that is equivalent to F203 (*SI Appendix*, Fig. S1).

Significance

Voltage-gated sodium (Na_v) channels control neuronal signaling and are key targets for local anesthetics, antiepileptics, and therapeutics for a range of disorders. Multimicrosecond Anton simulations have provided completely unbiased molecular-level views of the interactions of lipophilic drugs with the recently solved bacterial channel, Na_vAb from *Arcobacter butzleri*. Newly parameterized benzocaine and phenytoin molecules exhibit different membrane partition coefficients, crossing rates and distributions around the channel, leading to the identification of distinct high- and low-affinity sites. We observe a minimum free energy pathway through membrane-bound fenestrations to a pore-blocking location, or from aqueous solution directly through the (closed) intracellular gate. These observations help explain experimental data and provide insight into Na_v inhibition processes that will assist future drug development.

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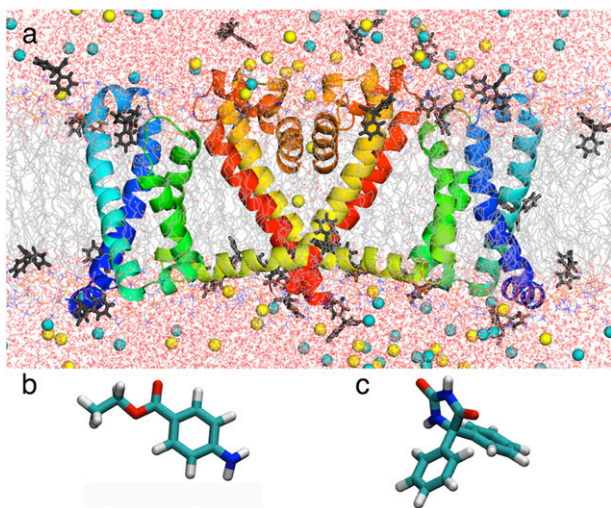


Fig. 1. (A) Simulation system showing Na_vAb (two of four subunits as ribbons, with voltage sensor domain (VSD) S1 and S2 in blue and S3 and S4 in green, pore domain (PD) S5 in yellow and S6 in red, P-loop P1, P2, and selectivity filter (SF; orange) in a hydrated lipid bilayer (chains as gray lines; water as red/white sticks; NaCl as yellow/cyan balls) and PHT (dark gray sticks). (B and C) Stick representations of BZC (B) and PHT (C).

In contrast, the other bacterial Na_v channels only possess two conserved Phe (F201 and F207 in Na_vAb), with orientations inconsistent with an FS6 site (participating in interactions away from the pore; Fig. 2B). Na_vAb may therefore offer unique similarity to mammalian Na_v for the investigation of LA/AE binding (we note, however, that binding may still occur in the absence of an FS6-like residue, as seen for brominated drug-like compounds to more distant F214 on S6 in Na_vMs; ref. 11).

In the 1970s, Hille proposed two separate pathways for inhibitor binding, with charged drugs such as titratable amines and quaternary ammonium derivatives requiring an open channel, and neutral drugs such as BZC and PHT entering via a lipophilic pathway somewhere in the membrane (allowing for binding and dissociation, even when the pore is nonconducting) (20). Our previous simulations revealed dynamic interplay between the protein and lipids, with fenestrations allowing lipid tails to enter the pore (21), representing a potential drug pathway (20, 22). However, access to these openings, and subsequent binding, have yet to be described. Here we use the Anton supercomputer (23) to carry out extensive unbiased fully atomistic simulations of BZC and PHT binding to the Na_vAb channel to reveal the distribution of binding sites and observe their pathways to shed light on inhibition mechanisms and aid future Na_v-drug development.

Results and Discussion

Membrane Partitioning of BZC and PHT. To examine the abilities of these amphiphilic drug molecules to move within the membrane, we first carried out Umbrella Sampling (US) simulations for a pure lipid bilayer, using newly parameterized standard CHARMM models (SI Appendix). SI Appendix, Fig. S24 shows that BZC binds at the membrane interface by -3.3 ± 0.4 kcal/mol, with center of mass (COM) at $|z| \sim 8.6 \pm 0.1$ Å from the bilayer center, interacting primarily with lipid carbonyls and phosphates via its amine H (SI Appendix, Fig. S34). The corresponding membrane partition coefficient for BZC, P_b , is 38.7 ± 6.3 , being less than the experimental values [ranging from 186 to 202 (24, 25) at 310–313 K]. This comparison with a range of results indicates ~ 1 kcal/mol error, and suggests somewhat too weak binding to the membrane interface, perhaps due to the slightly overestimated dipole moment (SI Appendix, Table SA4), chosen to ensure correct condensed phase properties (26, 27) and accurate interaction energies (SI Appendix, Table SA6). However,

SI Appendix, Table S1 shows that the model reproduces water–hydrocarbon partitioning free energies to within 0.2–0.3 kcal/mol. We consider these levels of discrepancy to be acceptable, being comparable to typical kcal/mol-order statistical errors, and expected to be largely cancelled for relative occupation of binding sites (see *Benzocaine Binding to the Channel*). BZC faces a smooth barrier of $+2.7 \pm 0.4$ kcal/mol (relative to interfacial minima; -0.6 ± 0.1 kcal/mol relative to water). Applying Kramer’s transition rate theory (SI Appendix), we obtain a crossing rate of $1.8 \pm 0.5 \mu\text{s}^{-1}$, indicating rapid submicrosecond membrane transfer.

PHT has a similar dipole moment to BZC (SI Appendix, Tables SA4 and SA5), yet experiences stronger binding at the membrane interface (SI Appendix, Fig. S24). The minimum of -4.1 ± 0.1 kcal/mol (at $|z| \sim 9.59 \pm 0.03$ Å), allows both aromatic rings to reside in hydrocarbon while satisfying H-bonds at the interface, to a greater extent than BZC (SI Appendix, Fig. S3B). There is a similar discrepancy between calculated (139 ± 25) and experimental (657 ± 33) (28) partition coefficients for PHT, although those experiments were done using a different membrane (egg phosphatidylcholine) and temperature (298 K). The partition coefficient is 3.6 times higher than that for BZC with our models, consistent with experiments (24, 25, 28). PHT experiences a high and sharp barrier of $+4.8 \pm 0.1$ kcal/mol relative to the interfacial minima ($+0.66 \pm 0.07$ kcal/mol with respect to water). We compute a crossing rate of $35 \pm 3 \text{ ms}^{-1}$ (SI Appendix); being ~ 2 orders of magnitude smaller than that for BZC, and approaching the millisecond time scale. This crossing rate suggests that PHT binding deep in the membrane may be restricted, even with multimicrosecond simulations.

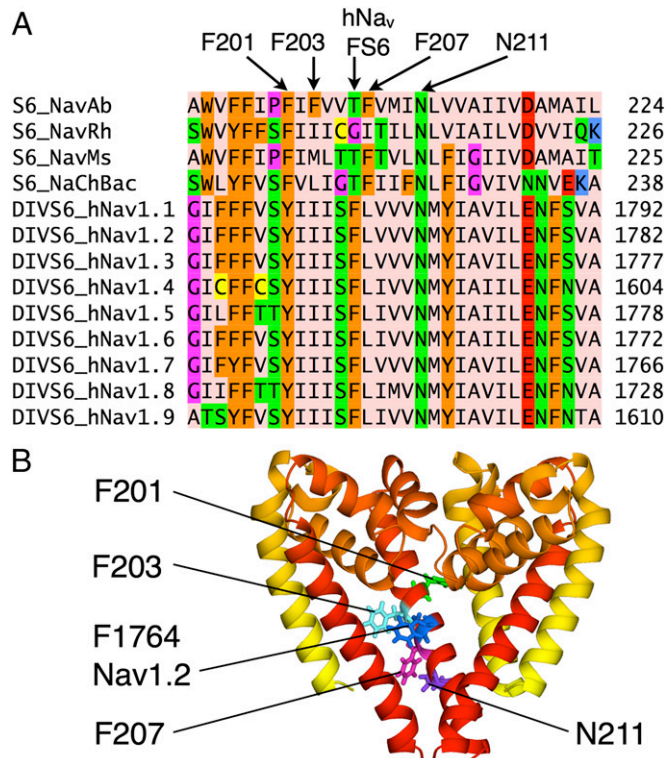


Fig. 2. (A) Sequence alignment of segment S6 of Na_vAb to other Na_v channels (showing only FS6-containing domain IV; see SI Appendix, Fig. S1, for full comparison). F201, F203, F207, and N211 in Na_vAb and F1764 in Na_v1.2 are indicated. Amino acids were colored with Jalview (46, 47) using the Zappo scheme. (B) Aligned structures (showing three subunits) of Na_vAb and model Na_v1.2 (based on Na_vAb) comparing Na_vAb F201, F203, and F207 to Na_v1.2 F1764. Conserved N211 (Na_vAb numbering) is also indicated.

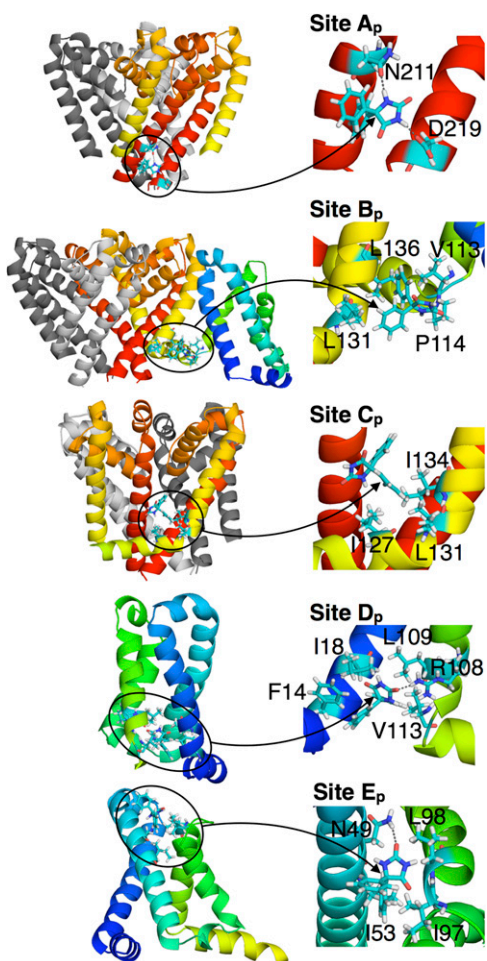


Fig. 5. Interactions involved in the binding sites A_p to F_p of PHT. (Insets) Arrows point directly at the PHT molecule.

BZC confinement in the central cavity and not due to direct stacking). Analysis of π -stacking conformations for outer and inner binding (*SI Appendix*, Fig. S6 *F* and *G*) shows that both favor more stable parallel displaced and T-shaped conformations (31). However, shorter distances are achieved from the outside (*SI Appendix*, Fig. S6 *E* and *F*), suggesting stronger interactions for initial binding. The ability of BZC to π -stack with F203 supports our assertion that F203 in Na_vAb plays a similar role to mammalian FS6, leading to a pore-blocking site.

Below this high-affinity site is B_b , deep in the intracellular gate, created by hydrophobic interactions with I217 and V213 (Figs. 3C and 4). Its K_D of $407 \pm 54 \mu\text{M}$ is higher (weaker) than A_b , and may be specific to the closed gate. Although both A_b and B_b evidence the ability for BZC to directly block the channel, the nonspecific binding in B_b may dominate closed-state low-affinity block, involving stabilization of the S6 bundle crossing via a hydrophobic seal (15).

Stacks of aromatics formed by pore helices create localized sites higher in the P-loop; this includes: C_b [intersubunit, F167(P1)–Y168(P1) and W195(S6), $K_D = 460 \pm 62 \mu\text{M}$], potentially linked to low-affinity binding (32); D_b , at the PD–voltage sensor domain (VSD) interface [F167(P1), Y142(S5) and F37(S1–S2), $K_D = 811 \pm 77 \mu\text{M}$]; and E_b [intrasubunit; F144(S5)–F198(S6)–F201(S6), $K_D = 4790 \pm 830 \mu\text{M}$]. Sites C_b and D_b , connected via F167, appear to be important in the access pathway (see *Drug Access Pathways*).

At a level just below fenestrations and between S5/S6 and the VSD, BZC binds to aromatic pocket site F_b , created by F107(S4), F140(S5), and F207(S6). This site may help explain block by voltage-sensor inhibition, suggested to occur close to FS6, restricting the bottom of the PD and S4–S5 linker, reducing

the effect of voltage on the activation gate (15, 33, 34). Even more directly influencing activation, low-affinity site G_b involves H-bonding to N49 and interaction via cation– π to the voltage-sensing R102 (S4; see Fig. 4). This finding is consistent with evidence for stabilization of the depolarized up state by LAs (33, 35).

Phenytoin Binding to the Channel. Unbiased $\sim 3\text{-}\mu\text{s}$ simulation has been used to sample PHT binding to Na_vAb , leading to a distribution (Fig. 3 *E* and *F*) that is very different to BZC. PHT entered the bilayer and protein within $\sim 3 \mu\text{s}$, but was unable to penetrate into the pore (although approached fenestrations on two occasions). This result may be explained by the greater binding to the membrane interface and slower time scales to enter the membrane core, calculated above. It is also consistent with experiments that demonstrate mutation of mammalian FS6 reduces PHT efficacy only marginally (36), suggesting other binding sites may play a greater role for this drug. Interestingly, the K_D for PHT sites A_p – E_p , (Figs. 3 *G* and *H* and 5) range from as little (as strong) as $19 \mu\text{M}$, to as high as $294 \mu\text{M}$ (*SI Appendix*, Table S3). Thus, the dominant PHT sites exhibit lower K_D values than seen for BZC, consistent with the higher affinity for PHT compared with BZC seen experimentally for mammalian channels (37). However, binding of PHT is of higher affinity than expected, given the closed state of the channel's activation gate [e.g., $K_D(\text{resting}) \sim 600 \mu\text{M}$ for mammalian Na_v (8)], perhaps indicating increased binding to non-FS6-like sites in Na_vAb due to sequence differences and fourfold symmetry, or due to increased interfacial preference for the PHT model used.

PHT site A_p is formed by residues in S6, involving H-bonding between imide H and conserved N211 side chain amide O, and D219 carboxyl O (conserved as D or E in mammalian channels) of adjacent subunits. A similar binding site has also been identified for a volatile general anesthetic in a NaChBac homology model, involving H-bonding to the equivalent residue N225 (14). Interestingly, mutation of N418 in DI, N1466 (DIII), and N1769 (DIV) in $\text{Na}_v1.2$, which aligns with N211 (Fig. 2 *A* and *B*), leads to reduced affinity for LAs (38). This site has previously been suggested to be pore-facing and to involve hydrophobic residues at the bottom of S6 (38). However, we observe that N211 forms A_p that is not pore facing in Na_vAb , and instead associate the nearby hydrophobic residues forming B_b with such a pore-facing site.

The binding of PHT at site A_p relies on the gate being in its closed, nonconducting state, locking the helices in place and effectively blocking the activation gate from both collapse, associated with slow inactivation (6, 21), and opening. *SI Appendix*, Fig. S7 shows that the presence of PHT in the simulation drastically modifies the behavior of the S6 helices at the level of the activation gate by limiting the natural fluctuations observed in the absence of drug (21) or in the presence of BZC. In particular, it can be seen that PHT reduces asymmetry of the gate (measured as the difference between the distances separating the COM of the C_α of residues 215–218 on pairs of opposite monomers), due to binding at A_p involving H-bonding to N211 and D219. The K_D for this site ($73 \mu\text{M}$) is consistent with the experimental value for binding to the NaChBac resting state ($120 \mu\text{M}$) (39).

Site B_p involves hydrophobic interactions with residues connecting the PD and VSD [L131(S5), L136(S5), V113(S4), and P114(S4–S5) of adjacent subunits, as well as Y20(S1)] and may contribute to low-affinity bundle crossing stabilization (37). Site C_p is the closest to the fenestrations (and F203), but is made up of nonspecific interactions with I127(S4–S5), L131(S5) and I134(S5). Site D_p is within the VSD and involves H-bonding of hydantoin carbonyl O to R108, with PHT phenyls interacting with F14(S1), I18(S1), L109(S4), and V113(S4). Similar site E_p involves imine H-bonding with N49(S2) and phenyl interactions with I53(S2), I97(S3), and L98(S4) (Fig. 5). *SI Appendix*, Table S3 summarizes the K_D values for PHT, revealing surprisingly high-affinity binding around the PD, VSD, and PD–VSD interface, without binding inside the pore. Again, this may explain the modest effect of FS6 mutation on PHT efficacy in mammalian channels (36).

Knowledge of the LA and AE binding sites has provided the necessary first step in understanding the molecular origins of Na_v channel inhibition. The next stage is to explore the state dependence of these interactions. Such studies are promising, given that F203 orientation and accessibility have been linked to structural changes associated with slow-inactivation (21), suggesting conformation-dependent drug binding within Na_vAb (10, 36, 40). Furthermore, by having uncovered a range of lower-affinity sites and the pathways of drug access, and not just the more conserved high-affinity blocking site, these results suggest new targets for therapeutically selective Na_v inhibitor compounds.

Methods

The Na_vAb protein (Protein Data Bank ID code 3RVY; ref. 3) was embedded in a bilayer of dipalmitoylphosphatidylcholine (DPPC) with explicit TIP3P water molecules and 150 mM of NaCl. High initial concentrations of 500 mM BZC (Fig. 1B) or 75 mM PHT (Fig. 1C; lower due to the tendency for PHT to aggregate in solution in trial μs -long trajectories) were added, with both systems experiencing rapid partitioning of drugs to the membrane, such that the membrane-channel system was in equilibrium with 5–10 mM bulk drug concentrations after $\sim 0.2 \mu\text{s}$ (SI Appendix, Fig. S4C). Systems were built and preequilibrated with CHARMM (41) and then further equilibrated using NAMD (42). Anton software from D. E. Shaw Research was then used for production runs, using the purpose-built Anton supercomputer (23) for additional 1.8 and 2.9 μs in the

presence of BZC and PHT, respectively. BZC and PHT topology and parameters were developed de novo, consistent with the CHARMM force field, with initial guesses for partial atomic charges and other force field parameters obtained using the CGENFF program (43, 44), optimized and validated using QM calculations and experimental partitioning free energies, as described in SI Appendix.

Free energy maps for drug binding to Na_vAb were calculated from unbiased Anton simulations as $W(R_i) = -k_B T \ln[\rho(R_i)] + C$, where ρ is the unbiased probability distribution as a function of reaction coordinates R_i (position vectors of the COM of the drug molecule relative to the COM of the PD, translated along z to align with the COM of the membrane), with C a constant. Binding sites were identified when lower than -1 kcal/mol in free energy relative to the membrane reference, with calculation of K_D values described in SI Appendix. Separate simulations of membrane partitioning were performed in dimyristoylphosphatidylcholine (DMPC) in 150 mM of NaCl. US (45) simulations for BZC and PHT were performed along the z axis with the drug COM held with respect to membrane COM near each US window center by a $2.5 \text{ kcal/mol}/\text{\AA}^2$ force constant for 13–25 ns each. Full details of US simulations, partition coefficient, and rate calculations are provided in SI Appendix.

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