Tetrameric Bacterial Sodium Channels: Characterization of Structure, Stability, and Drug Binding†

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ABSTRACT: NaChBac from Bacillus halodurans is a bacterial homologue of mammalian voltage-gated sodium channels. It has been proposed that a NaChBac monomer corresponds to a single domain of the mammalian sodium channel and that, like potassium channels, four monomers form a tetrameric channel. However, to date, although NaChBac has been well-characterized for functional properties by electrophysiological measurements on protein expressed in tissue culture, little information about its structural properties exists because of the difficulties in expressing the protein in large quantities. In this study, we present studies on the overexpression of NaChBac in Escherichia coli, purification of the functional detergent-solubilized channel, its identification as a tetramer, and characterization of its secondary structure, drug binding, and thermal stability. These studies are correlated with a model produced for the protein and provide new insights into the structure—function relationships of this sodium channel.

Voltage-gated sodium channels are integral membrane proteins that selectively transport sodium ions across cellular membranes in response to changes in membrane potential. Their function in mammals underlies neuronal signaling, muscle contraction, and hormone secretion (1). The first bacterial voltage-gated sodium channel (NaChBac) was identified from Bacillus halodurans (2), and a number of orthologs were subsequently identified, forming a superfam-

ily (3). The function of voltage-gated sodium channels in bacteria is still a matter of debate; however, there is evidence that these channels are involved in processes such as motility, chemotaxis, and pH homeostasis (4).

The pore-forming α-subunit of mammalian voltage-gated sodium channels consists of a single polypeptide of 24 transmembrane α-helices arranged as a pseudotetramer of four homologous domains (designated DI—DIV), with each domain comprised of six transmembrane (6TM) helices (designated S1—S6). In contrast, NaChBac appears to consist of a single 6TM domain of 274 amino acids, and it has been proposed to form a functional unit as a tetramer (1), although to date there has been no direct structural evidence to demonstrate that. Its kinetics of activation, inactivation, and recovery from inactivation are 10−100 times slower than those of mammalian voltage-gated sodium channels, and although NaChBac is sodium selective, its high degree of homology to calcium channels in the pore domain allows it to be converted into a calcium selective channel by three aspartic acid point mutations (5). In addition, NaChBac can be blocked by calcium channel blockers (2). Further func-
tional studies of NaChBac combined with mutagenesis have identified important residues involved in channel activation (6–8) and inactivation (9). The identification of glycine 219 as a critical residue involved in channel activation has shed light on what may be a general mechanism for voltage-gated ion channel gating (6).

Sodium channels, like other membrane proteins, are notoriously difficult to study in vitro due to their generally low natural abundance in vivo and the difficulties with overexpression, detergent solubilization, and purification in an active form. Previous structural studies on sodium channels have consequently tended to focus on channels from electric eels, in which they are naturally present in high abundance in electroplax tissue (10–12). To date, expression of NaChBac has been reported in only cell-based systems for electrophysiological studies, and for luminescence studies, which only require small amounts of protein (13).

Information about NaChBac is important for our understanding of structural and functional properties that are common to both bacterial and eukaryotic channels, not only because of their sequence homology but also because NaChBac shares functional features common to all three eukaryotic voltage-gated ion channel families (14). In this paper, we report on the successful expression and purification of milligram quantities of folded functional protein and have carried out a series of structural and functional biophysical characterizations, establishing its secondary structure, oligomeric state, drug binding, and thermal stability.
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EXPERIMENTAL PROCEDURES

Materials. The cDNA of NaChBac from *B. halodurans* was generously provided by D. Clapham of Harvard Medical School (Boston, MA). *Escherichia coli* lipids were purchased from Avanti Polar Lipids, while the detergent dodecyl maltoside (DDM) was obtained from Anatrace. Mibefradil dihydrochloride hydrate was from Sigma Aldrich, and sodium green dye was from Invitrogen.

Cloning. Full-length NaChBac cDNA (274 amino acids) was amplified by PCR using *Pfu* DNA polymerase (Stratagene) and two oligos: F, 5′-GCCATATGGAAATG-GAAGCTAGACAG-3′ with an NdeI site at the 5′ end; and R, 5′-CGGATCCCTATTACGGATTTAAGC-3′ with two stop codons followed by a BamHI site at the 5′ end. The PCR product recovered from a 1% agarose gel was digested with NdeI and BamHI and inserted into a pET15b vector (Novagen) between NdeI and BamHI sites using T4 DNA ligase. The ligation mixture (2 μL) was used for transformation into DH5α cells. Once the transformation process was complete, the cells were spread on agar plates containing 100 μg/mL ampicillin and incubated at 37 °C overnight. Five colonies were picked from the plates, and DNA was prepared (pET15b/NaChBac). All colonies were confirmed by sequencing to contain a (His)6 tag followed by a thrombin digestion site at the N-terminus of the wild-type NaChBac gene.

Expression. Competent cells, the DE03 C41 strain of *E. coli*, were transformed by pET15b/NaChBac, and five colonies from the overnight agar plate were used to test induction. The colonies were grown in LB medium (Merck) with 100 μg/mL ampicillin and incubated at 37 °C overnight. Five colonies were picked from the plates, and DNA was prepared (pET15b/NaChBac). All colonies were confirmed by sequencing to contain a (His)_6 tag followed by a thrombin digestion site at the N-terminus of the wild-type NaChBac gene.

Purification of NaChBac. Two flasks each containing 500 mL of LB+AMP were inoculated with 5 mL of overnight culture and the cells grown at 37 °C for 3 h. After cooling to 22 °C and being induced with IPTG, the cells were incubated for a further 72 h. The cells were collected and resuspended in 50 mM Tris (pH 7.7) and 500 mM NaCl (buffer A) and left at −80 °C overnight. The cells were thawed at room temperature and sonicated on ice for 2 × 10 min in the presence of DNAase I (2 μg/mL), lysozyme (2 mg/mL), and 0.2 mM PMSF (VWR International). Disrupted cells were centrifuged at 10000g for 30 min. The resulting pellet was discarded and the supernatant centrifuged at 250000g for 2 h. The pellet, which mainly contains membranes, was resuspended in 180 mL of buffer A, and DDM was added to a final concentration of 1% and incubated overnight at 4 °C. Insoluble materials were removed by high-speed centrifugation, and after 20 mM imidazole was added, the supernatant was loaded on a 5 mL His Trap nickel column (GE Healthcare) that was equilibrated with buffer A having 20 mM imidazole (and 1% DDM), at a flow rate of 0.5 mL/min. After washing steps with buffer B [500 mM NaCl, 50 mM Tris-HCl (pH 7.7), and 0.1% DDM] containing imidazole (first with 50 mM and then with 100 mM), NaChBac was eluted from the column with 500 mM imidazole in buffer B.

Elution fractions were concentrated in a Amicon concentrator (100 kDa cutoff). Some impurities (proteins smaller than ~100 kDa) passed through the concentrator during this step.

Samples (2 mL) of the concentrated material were loaded on the top of a gel filtration column (Superdex200, Bio-Rad) in buffer B at 4 °C and eluted at a rate of 0.5 mL/min. The purity of fractions was assayed by SDS gel electrophoresis, and the identity of the single dominant band was confirmed by mass spectroscopic analysis.

For Western blotting, fractions from the gel filtration column were run on a precast denaturing gel (NuPAGE 4 to 16% gradient, Invitrogen) and transferred to a polyvinyl difluoride (PVDF) membrane. Anti-His G/A/P antibody (Invitrogen) was used as the primary antibody, and goat anti-mouse IgG AP conjugate (Novagen) was used as the secondary antibody. Immunoblotting signals were developed with BCIP/NBT FAST tablets (Sigma).

Blue Native Gel. To examine the oligomeric state of native NaChBac in solution, fractions from the gel filtration column were run on a 4 to 16% gradient Native PAGE Bis-Tris gel (Invitrogen) with 1% DDM at 4 °C. NativeMarker Unstained Protein Standards, 20–1230 kDa (Invitrogen), were used. To dissociate the oligomers, some NaChBac samples were treated with 1% SDS prior to being applied to the gel.

Dynamic Light Scattering. Dynamic light scattering recordings were made using a DynaPro-801 instrument. The change in diffusion coefficient as a function of protein concentration was measured, and extrapolation of the diffusion coefficient at “zero” concentration was used to calculate the sample molecular mass, using a standard curve of known diffusion coefficients and molecular masses. Initial protein concentrations of approximately 2 mg/mL were used, and for subsequent measurements, a series of three 1:1 dilutions were made using phosphate-buffered saline containing 0.1% DDM. Measurements were analyzed in bimodal mode, and an average of at least 25 measurements was used at each concentration.

Circular Dichroism (CD) Spectroscopy. CD spectra were recorded using an Aviv 215 spectropolarimeter equipped with a detector acceptance angle of >90° (for membrane/scattering samples) that had been calibrated using camphorsulfonic acid for optical rotation and benzene vapor for wavelength. Spectra were collected at 0.2 nm intervals over the wavelength range from 280 to 185 nm. Three scans were collected for each protein sample and its corresponding baseline (which contained buffer and 0.1% DDM) in the same 0.01 cm path length Suprasil cuvette. Measurements were only made down to wavelengths at which the instrument dynode voltage indicated that the intensity of the light reaching the detector was sufficient for accurate detection (15). Using the CDtools processing package (16), the averaged baseline spectra were subtracted from the corresponding averaged protein spectra, and the spectra were smoothed with a Savitsky-Golay filter.
Spectra were analyzed for secondary structure using the Dichroweb server (17) with the SP175 reference data set (18) and the CDSSTR algorithm (19), using a mean residue weight value of 114.8.

For thermal scans, measurements were made on samples in DDM (0.95 mg/mL protein) at 222 nm using intervals of 2 °C; an equilibration time of 5 min was used at each temperature point. For the samples containing mibefradil, a final drug concentration of 200 mM was used.

**Flux Assay.** We prepared lipid vesicles by dissolving *E. coli* lipids in a 1:1 chloroform/methanol mixture and freeze-drying them under vacuum. Lipids were then rehydrated to 20 mg/mL lipid in 1 mL of 20 mM Tris (pH 7.5) containing 10 µM sodium green dye (20). Liposomes with a diameter of ∼100 nm were formed by extrusion using a mini extruder (Avanti Polar Lipids) (21).

NaChBac reconstitution was achieved by rapid dilution of DDM-solubilized protein into liposomes, producing a final lipid:protein weight ratio of 100:1. Initially, 500 µL of 20 mg/mL liposomes was added to 100 µg of NaChBac (3.4 mg/mL in 0.1% DDM) and sonicated for 10 s. Subsequently, DDM was diluted to below its critical micelle concentration by a 10-fold dilution of the sample with 5 mL of 20 mM Tris buffer (pH 7.5) and continuous mixing at room temperature for 20 min. Proteoliposomes were pelleted by centrifugation at 650000g and 25 °C for 60 min. Finally, proteoliposomes were resuspended in 1 mL of Tris buffer (pH 7.5).

Fluorescence flux experiments were carried out using a FluoroMax-3 spectrofluorometer (Horiba Jobin Yvon Ltd.) at 25 °C. Uptake of sodium into proteoliposomes was initiated by a 33-fold dilution of liposomes containing NaChBac into 20 mM Tris buffer (pH 7.5) containing 300 mM sodium chloride. Changes in sodium green fluorescence were monitored at 540 nm (1 nm bandwidth, excitation at 512 nm) following a mixing time of ∼10 s, after which the increase in fluorescence had stabilized. NaChBac activity was assayed both in the presence and in the absence of the channel blocker mibefradil (2). Mibefradil (200 µM) was added to the proteoliposome sample and incubated for 5 min at room temperature prior to the fluorescence experiments. Following the same procedure that is described above, control experiments were carried out in the absence of protein (liposomes without NaChBac but still in the presence of DDM), in the presence of mibefradil without protein (to test its effect on leakage), and in the absence of sodium chloride by diluting proteoliposomes or liposomes into sodium-free buffer. Each experiment was repeated three times with the exception of the controls carried out in the presence of sodium chloride-free buffer, which were repeated twice. As a final test of the maximal fluorescence response, NaChBac-containing liposomes were also treated by incubating with 2% DDM at room temperature for 10 minutes to break up the liposomes, and the response was recorded.

**Molecular Modeling.** A homology model of the NaChBac transmembrane regions was generated using the crystal structure of the chimeric Kv1.2-Kv2.1 potassium channel (22), as this structure offers a complete set of coordinates for the voltage sensor region. Although the choice of structural template differed, our sequence alignment (Figure 4a) closely matched that of Blanchet et al. (8, 23); T-COFFEE (24) was used to generate the sequence alignment, with the first positive charge of the NaChBac S4 helix aligned with the “R2” position of the potassium channel S4 helix. The accuracy of this alignment is supported by mutagenesis and functional studies on mutant NaChBac channels with substituted S4 charged residues (25).

Residues of the chimeric Kv1.2-Kv2.1 potassium channel structure (Protein Data Bank entry 2R9R) were mutated to the NaChBac sequence using the Biopolymer module of SYBYL (version 7.0, Tripos Inc., St. Louis, MO). The model was subjected to 500 rounds of conjugate-gradient minimization in SYBYL using the Tripos force field (23). The model figure was produced using PyMOL (DeLano Scientific, San Carlos, CA). The number of residues in helical conformations in the final model was identified using the DSSP algorithm (26). Prediction of the secondary structure for NaChBac regions not encompassed by the model (including the S5–S6 extracellular linker that forms the selectivity filter) was conducted according to the consensus of a number of secondary structure prediction algorithms (27).

**RESULTS**

**Expression and Purification.** Initial attempts to express NaChBac in *E. coli* were fraught with difficulties (28) and resulted in either cell death, inclusion bodies, or no expressed protein. To achieve successful overexpression in sufficient quantities for biophysical characterizations, a number of modifications were required, including change of plasmid, moving the (His)_6 tag from the C-terminus to the N-terminus, growth in complex medium, and growth at low temperatures. Eventually, conditions were found (as described in this paper) in which the induced expression level for NaChBac (Figure 1) was quite high relative to the level of expression of other membrane proteins in bacterial systems. Furthermore, under these conditions, the background (bacterial protein expression) was partially inhibited, which resulted in both a high yield and fewer impurities being present prior to purification. After a one-step purification on a nickel column, the yield was roughly 3 mg/L of culture and the impurities made up <10%, as evidenced by gel electrophoresis (Figure 2a). The identity of the main band was shown by mass spectrometry.
to be bona fide; the identity of the only other significant band (slightly lower molecular mass) was also shown to be derived from NaChBac. It appeared to be a large C-terminal fragment (consistent with it missing the N-terminal end due to its lack of staining with antibodies against His tags on the Western blots) (data not shown). Because it was retained on the nickel column with the His-tagged proteins, we suggest it may have been a proteolyzed monomer (that was retained as part of tetrameric complex, in which at least one monomer had a His tag enabling it to be retained on the column). Using a concentrator having a 100 kDa cutoff, impurities of less than ∼100 kDa were effectively removed. Following a further purification step on a gel filtration column, intact NaChBac was seen to be effectively the only band that could be detected on a SDS gel (Figure 2b), and on Western blots (Figure 2c). The total yield of purified protein was ∼2 mg/L of culture. To examine the stability of the protein with time, a SDS gel and Western blot were obtained with the purified protein that had been maintained at 4°C for 2 weeks. They were indistinguishable from those of the initial sample, exhibiting no detectable degradation products (data not shown).

**Oligomeric State.** To estimate the molecular mass and, therefore, the oligomeric state of the isolated protein in DDM, we ran a gel filtration column with the purified protein and found a single major peak. However, because the molecular mass calibration curve was made by running soluble standard proteins on the column, the estimated molecular mass was not expected to be highly accurate for a membrane protein that will have hydrodynamic properties different from those of soluble proteins. Nevertheless, the resulting molecular mass was estimated to be ∼227 kDa. Since a tetrameric NaChBac protein would be around 120 kDa and the contribution from the detergent in the DDM micelles should be ∼80−100 kDa, it is reasonable to assume that the protein complex isolated is the tetrameric form of NaChBac. Minor, later retained fractions corresponded to a molecular mass of ∼112 kDa, which would be compatible with a monomeric NaChBac (30 kDa) since the DDM micelles contribute 80−100 kDa.

Dynamic light scattering studies were also conducted to assess the molecular mass of the DDM-solubilized component. The sum of squares (SOS) fitting error which is defined as the sum of squares difference between the predicted and measured autocorrelation functions was consistently <2, indicating a good fit to the predicted autocorrelation function which consists of a single exponential. Large SOS values would indicate the experimental curve is multimodal and/or noisy. The measurements also showed that approximately 98% of the sample by mass was in a single state. The estimated molecular mass was 245 kDa, reasonably similar to the value obtained by gel filtration, again with some uncertainty due to the contribution of the detergent, but most consistent with a tetrameric complex with detergent.

To confirm the oligomeric nature, the DDM-solubilized material was run on a Blue native gel (Figure 2d). The protein migrated as a sharp peak with a total molecular mass of ∼220 kDa, again indicative of a tetrameric state of NaChBac in a mixed protein/detergent micelle and consistent with the gel filtration and dynamic light scattering results. The small amount of higher-molecular mass forms likely consists of aggregates that are common when membrane proteins are run on native gels. Finally, for comparison, the DDM-solubilized protein was treated with SDS (to dissociate it into monomers) prior to being run on a native gel (Figure 2d).

**Secondary Structure and Thermal Stability.** The CD spectrum of NaChBac in DDM (Figure 3a) is consistent with a protein that has a high helical content, having negative peaks at ∼222 and 208 nm and a large positive peak at ∼195
nm. Its spectrum is similar in shape to that of the purified sodium channel isolated from *Electrophorus electricus* in DDM (Figure 3a), except that it has a slightly larger magnitude and a red-shifted positive peak, both indicating it has a higher overall helix content. This is consistent with the longer expected loops and termini of the eukaryotic protein, which, if partially nonhelical, would result in an overall lower percentage of helix. This similarity to eel protein in both DDM and genapol (11) detergents is heartening because the eel channels have been shown to be both properly folded and functional (11; K. Charalambous et al., manuscript in preparation).

The secondary structure of NaChBac, as determined from analyses of its CD spectra, is 67% helix, 8% sheet, and 25% other. The goodness-of-fit parameter (NRMSD) of 0.010 suggests that the calculated secondary structure corresponds well with the experimental data. This is consistent with the molecular model (Figure 4b) for the transmembrane domains of the protein: the helical sections of this model comprise 45% of the total residues. Additional helices predicted in the extramembranous regions (data not shown) would increase the total helix content to one similar to that calculated from the CD data, i.e., approximately 70%. The calculated secondary structure of the eel channel from its CD spectrum was slightly lower (50%) (11), which is again consistent with the relative appearances of its spectrum and the spectrum of NaChBac, and with the observation that it has longer loops and termini that will have lower helical contents.

Upon addition of the channel-blocking drug mibefradil, the spectrum of NaChBac changes slightly, with a decrease in the magnitude of the 195 nm peak, but there is little change in the rest of the spectrum (Figure 3b). This might suggest that the drug not only binds to NaChBac in DDM micelles but also causes a small conformational change, resulting in a slight decrease in helix content.

The thermal stability of NaChBac in DDM was assessed using CD spectroscopy by monitoring the decrease in the magnitude of the peak at 222 nm (Figure 3c), which is associated primarily with helix content. The protein spectrum exhibited a characteristic cooperative two-state unfolding process, which was also as could be expected for a properly folded protein. Surprisingly, the protein appeared to be very stable, with an estimated $T_m$ of $\sim 60^\circ$ C. This was similar to the $T_m$ for the eel channel in the same detergent (Figure 3c). Interestingly, however, the total loss of helix content appears to be larger for the NaChBac protein. Even after “unfolding”, both proteins retained a significantly helical structure. In addition, the presence of mibefradil during unfolding (data not shown) increased the stability of the NaChBac protein even further, which is often the case when ligands bind to proteins in their active conformations.

Finally, the CD spectra of NaChBac in DDM (where it is a tetramer) and in SDS (where the tetramer dissociates into a monomer) were compared (Figure 3d). The monomer, like the thermally unfolded protein, still retains a significant helical structure, again suggesting the secondary structure of this protein is significantly stable but also that the association into tetramers results in secondary structural changes that make the protein functionally competent.

**Flux Activity.** To demonstrate that the tetrameric protein is functional, a flux assay in the presence and absence of the channel blocker mibefradil was undertaken. The assay is based on quantitation of the increase in the fluorescence of the dye sodium green in the presence of sodium. When sodium enters the vesicles containing the dye, the fluores-

![Figure 3: Circular dichroism spectra of NaChBac in DDM.](image-url)
cence increase can be monitored at 540 nm. The control samples consisted of proteoliposome samples in which the added buffer did not contain sodium, and experienced only a very small increase (<10% of the maximal increase observed following addition of sodium to proteoliposome samples). Liposomes alone (without protein) did experience an increase due to leakage, but this increase was significantly smaller than the increase in the proteoliposome samples which represented channel-induced flux (Figure 5). As a further control, proteoliposomes were treated with a channel blocker, mibefradil; the resulting fluorescence in those samples was not greatly different from that of the liposomes alone, suggesting that channel flux was indeed responsible for the increase in fluorescence, and that it could be prevented by blocking the channel. The presence of mibefradil itself (in liposomes without protein) did not significantly change the level of leakiness. As a final control, channels blocked by mibefradil but without sodium present exhibited fluorescence indistinguishable from that of the channels without sodium (and hence the change in fluorescence is not related to a spectroscopic property of the drug). The maximal level of the fluorescence response (for NaChBac-containing liposomes that were subsequently dissolved in detergent) demonstrated a small but significant further increase, suggesting that not all the liposomes necessarily contained active

Figure 4: Homology model of NaChBac transmembrane segments. (a) Sequence alignment of the TM segments (designated S1–S6) of the Kv1.2-Kv2.1 chimera (top row) and NaChBac (middle row). The bottom row indicates identical residues (asterisk), conserved residues (colon), and semiconserved residues (period). (b) Side view (perpendicular to the membrane plane) of a ribbon depiction of the model of the TM regions showing the four monomers present in the tetramer in different colors.
protein, or that the proteins present were not all oriented in the same direction in the reconstituted liposomes.

These studies have thus demonstrated that the purified NaChBac protein can be successfully reconstituted into liposomes, producing functional channels that not only transport sodium but are also blocked by a known channel blocker.

**DISCUSSION**

The overexpression and purification of membrane proteins are much more challenging than those of soluble proteins (29, 30). In particular, overexpression of membrane proteins can often cause cell death, which was the case in many of our NaChBac expression trials. This may be due to interference with components of the membrane essential for normal cellular functions and the necessity for maintenance of the intracellular ion content. In addition, protein yields tend to be limited by the finite capacity of the cell membrane system, and the ability to fold properly into the host membrane. In addition to difficulties in expression, extraction of protein from the membrane in a functional state can also be problematic.

We have developed a method for overexpressing the bacterial sodium channel from *B. halodurans* in *E. coli* in a functional form that enabled a wide range of biochemical and biophysical characterizations of this protein to be carried out.

It had been proposed that the NaChBac channel would be composed of four monomers forming a tetrameric assembly, similar to that found in both bacterial and eukaryotic potassium channels, but unlike mammalian sodium channels, which are a single polypeptide chain forming a pseudotetramer of four homologous domains. This study has demonstrated for the first time that the functional oligomeric state of NaChBac is indeed a tetramer. Furthermore, CD spectroscopic studies have shown that the measured secondary structure is consistent with what is expected for a 6TM channel based on homology modeling.

The DDM-solubilized protein is not only folded and tetrameric but also capable of binding the channel blocker mibebradil, as shown by both the flux assay and the CD binding study. The protein shows considerable stability to both thermal denaturation and treatment with SDS, which dissociates the monomers, both of which result in some loss of helix content but retention of a largely helical polypeptide. Such stability of a membrane protein helical core is reminiscent of bacteriorhodopsin (31), which also remains mostly helical even after unfolding in SDS, and KcsA, which even retains its tetrameric structure in SDS (32). The resistance to thermal stress in the anisotropic environment of a detergent micelle may reflect its requirement for flexibility, enabling it to undergo significant conformational changes associated with state-dependent functional changes.

In conclusion, we have demonstrated we can isolate and purify an active sodium channel protein and have characterized its secondary and quaternary structures, its thermal stability, and its binding of the drug mibebradil.

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