Molecular Architecture of the Voltage-dependent Na Channel: Functional Evidence for α Helices in the Pore

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ABSTRACT
The permeation pathway of the Na channel is formed by asymmetric loops (P segments) contributed by each of the four domains of the protein. In contrast to the analogous region of K channels, previously we (Yamagishi, T., M. Janecki, E. Marban, and G. Tomaselli, 1997. *Biophys. J.* 73:195–204) have shown that the P segments do not span the selectivity region, that is, they are accessible only from the extracellular surface. The portion of the P-segment NH2-terminal to the selectivity region is referred to as SS1. To explore further the topology and functional role of the SS1 region, 40 amino acids NH2-terminal to the selectivity ring (10 in each of the P segments) of the rat skeletal muscle Na channel were substituted by cysteine and expressed in tsA-201 cells. Selected mutants in each domain could be blocked with high affinity by externally applied Cd2+ and were resistant to tetrodotoxin as compared with the wild-type channel. None of the externally applied sulfhydryl-specific methanethiosulfonate reagents modified the current through any of the mutant channels. Both R395C and R750C altered ionic selectivity, producing significant increases in K+ and NH4+ currents. The pattern of side chain accessibility is consistent with a pore helix like that observed in the crystal structure of the bacterial K channel, KcsA. Structure prediction of the Na channel using the program PHDhtm suggests an α helix in the SS1 region of each domain channel. We conclude that each of the P segments undergoes a hairpin turn in the permeation pathway, such that amino acids on both sides of the putative selectivity filter line the outer mouth of the pore. Evolutionary conservation of the pore helix motif from bacterial K channels to mammalian Na channels identifies this structure as a critical feature in the architecture of ion selective pores.

KEY WORDS: pore helix • cysteine mutagenesis • tetrodotoxin • methanethiosulfonates • cadmium

INTRODUCTION

In voltage-gated ion channels the pore-lining (P) segments in each of four internally homologous domains (Na and Ca channels) or subunits (K channels) have been proposed to form hairpins that partially traverse the membrane and together create an ion-selective pore (Stuhmer, 1991; Catterall, 1995; Marban et al., 1998). Based on mutagenesis data and homology with a ring of glutamic acid residues in the Ca channel, the sequence-aligned residues in the Na channel pore D400, E755, K1237, and A1529, (DEKA; see Fig. 1) form at least part of the selectivity filter (Heinemann et al., 1992). The external mouth of the pore has been extensively studied by site-directed mutagenesis. Mutations introduced on the COOH-terminal side of the DEKA ring have been electrophysiologically characterized using toxin or metal cation blockers as structural probes (Backx et al., 1992; Satin et al., 1992; Dudley et al., 1995; Favre et al., 1995, 1996; Chiamvimonvat et al., 1996; Perez-Garcia et al., 1996; Chen et al., 1997; Li et al., 1997, 2000a,b; Tsushima et al., 1997a,b; Yamagishi et al., 1997; Chang et al., 1998; Penzotti et al., 1998; Sunami et al., 2000). High affinity Cd2+ block and modification by extracellular methanethiosulfonate (MTS) reagents of cysteine substitution mutants on the COOH-terminal side of the DEKA ring suggests that this part of the pore is composed of random coils rather than periodic structures such as α helices or β strands (Chiamvimonvat et al., 1996; Perez-Garcia et al., 1996; Tsushima et al., 1997a,b; Yamagishi et al., 1997). The topology of the amino acid residues immediately NH2-terminal to the DEKA ring (Yamagishi et al., 1997) is distinct from that in voltage-dependent K channels (MacKinnon and Yellen, 1990; Yellen et al., 1991). Cysteine mutants on the NH2-terminal side of the DEKA ring are not accessible from the internal mouth of the channel; in fact, two substitutions immediately NH2-terminal of the DEKA residues (F1236C and T1528C) are accessible from the external surface (Yamagishi et al., 1997). It appears that the P segments of the Na channel do not traverse the membrane as they do in K channels (MacKinnon and Yellen, 1990; Yellen et al., 1991).

As might be predicted by the sequence divergence of the P segments in each domain of the Na channel, the
contribution of each of the segments to the formation of the pore is not equivalent (Chiamvimonvat et al., 1996). Amino acids in each of the P segments contribute differently to the permeation phenotype of the channel; residues in the DEKA ring importantly influence conductance (DEK) and ionic selectivity (K; Heinemann et al., 1994; Chiamvimonvat et al., 1996; Favre et al., 1996; Perez-Garcia et al., 1997; Li et al., 2006). Other residues in the domain IV P segment (Gl530, W1531, and D1532) are major determinants of monovalent cation selectivity (Chiamvimonvat et al., 1996; Tsuchiya et al., 1997b).

The crystal structure of the pore of a bacterial K channel (KcsA) exhibits several features predicted to be conserved across the superfamily of voltage-gated ion channels (Doyle et al., 1998). A pore helix is present immediately NH2-terminal to the selectivity filter of KcsA. The pore helix is a critical determinant of the structural integrity of the pore and ensures the proper alignment of the selectivity filter residues. However, Na channels must be structurally distinct from K channels based on the limited amino acid homology and different ion selectivity. It is uncertain which, if any, of the structural motifs in KcsA have been retained in the Na channel pore. Structural modeling suggests that the P segments of the Na (Hsu et al., 1999; Lipkind and Fozzard, 2000) and Ca channels (Hsu et al., 1999) retain the α-helix-turn-coil motif described in the KcsA structure (Doyle et al., 1998). We sought to test experimentally the structure predictions of the P segments of the Na channel using a combination of cysteine mutagenesis, electrophysiological recording, and site-specific modification. Cysteine substitutions were made at 40 positions NH2-terminal to the DEKA residues in the α1-skeletal muscle Na channel and expressed in tsA-201 cells. Cd2+ and tetrodotoxin (TTX) block or MTS modification gauged the external accessibility of these mutants, and monovalent cation selectivity was examined by ionic substitution. The experimental data are incorporated into a revised structural model of the Na channel P segments.

MATERIALS AND METHODS

Mutagenesis and Channel Expression

Mutagenesis of the expression plasmid containing the rat skeletal muscle (μ1) α subunit of the Na channel was performed as described previously (Yamagishi et al., 1997). The wild-type μ1 Na channel α subunit cDNA (Trimmer et al., 1989) was cloned into the mammalian expression vector pGWIH for both mutagenesis and expression. The rat brain β1 subunit (Isom et al., 1996) was cloned into pCMV-5 (Phillipson et al., 1993). A plasmid expressing the green fluorescent protein (pGreenLantern; Gibco BRL) was used as a transfection reporter. Cysteine substitutions were introduced into the μ1 cDNA by oligonucleotide-directed mutagenesis (Perez-Garcia et al., 1996) or by PCR with overlapping mutagenic primers (Quickchange™ site-directed mutagenesis kit; Stratagene). All mutations were performed in duplicate and confirmed by DNA sequencing the region of the mutation. TsA-201 cells, a transformed human embryonic kidney cell line (HEK 293) stably expressing the SV40 T-antigen, were used for channel expression. The culture and transfection conditions were as described previously (Yamagishi et al., 1997). Transfection was performed by the calcium phosphate precipitation method for 6–12 h with a combination of 2.5 μg of the wild-type or mutant α subunit, 1 μg of the β1 subunit, and 0.3 μg of the green fluorescent protein cDNA-containing plasmids per 35-mm dish.

Electrophysiology and Data Analysis

Transfected cells were identified by epifluorescence microscopy and membrane currents were recorded with the whole-cell configuration of patch clamp (Hamill et al., 1981) 24–72 h after transfection. Cells were transferred to the stage of an inverted microscope and superfused with external solution at a rate of 1–2 ml/min. All experiments were performed at room temperature (22–23°C). Patch electrodes were pulled from borosilicate glass and had 2–5 MΩ tip resistances. Currents were recorded using a patch-clamp amplifier (model Axopatch 200A; Axon Instruments) interfaced to a personal computer. Cell capacitance was calculated by integrating the area under an uncompensated capacity transient elicited by a 20-mV hyperpolarizing test pulse from a holding potential of ~80 mV. Series resistance was then compensated as much as possible without ringing, typically 70–90%. Given the average series resistance of our electrodes, the maximal uncompensated voltage error was <6 mV for the largest currents studied. Whole-cell currents were recorded in a bath solution containing (in mM): 140 NaCl, 5 KCl, 2 CaCl2, 1 MgCl2, 10 HEPES, and 10 glucose, pH 7.4. The pipette solution was (in mM): 35 NaCl, 105 CsF, 1 MgCl2, 10 HEPES, and 1 EGTA, pH 7.2. Dose–response curves for block of the mutant channels by extracellular Cd2+ were determined by adding the chloride salt of the blocker to the bath at concentrations between 1 μM and 5 mM, and TTX was added at concentrations between 1 nM and 10 μM, depending on the sensitivity of the particular mutant. Full current-voltage relationships were determined at each Cd2+ or TTX concentration. The half-blocking concentration (IC50) was determined by a least-squares fit (Levenberg-Marquardt algorithm; Microcal Origin) of the data to the function: I/I0 = 1/[1 + (blocker)/IC50], where I0 and I are the currents in the presence and absence of blocker, respectively. In nearly all cases, block by Cd2+ and TTX was reversible, when the current amplitude did not return to >90% of control channel rundown could not be excluded and the experiments were not included in the analysis.

Susceptibilities to extracellular methanethiosulfonate ethylammonium (MTSEA), methanethiosulfonate ethyltrimethylammonium (MTSET), methanethiosulfonate ethylsulfonate (MTSES; Toronto Research Chemicals) modification was determined by bath application of saturating concentrations of each reagent (2.5 mM MTSEA, 1 mM MTSET, and 5 mM MTSES). MTS modification of the current was verified by the irreversibility of the change in the current amplitude. In no case was there an irreversible change in current amplitude after exposure to MTS reagents.

Single-channel recording was performed in those mutant channels that exhibited sensitivity to block by Cd2+ at the whole-cell level. The cell-attached configuration of the patch clamp was used (Hamill et al., 1981). The currents were sampled at 10 kHz and low-pass filtered at 2 kHz. The pipette solution contained (in mM): 140 NaCl, 10 HEPES, pH 7.4, and 400 μM Cd2+. The bath contained (in mM): 140 KCl, 1 BaCl2, 10 HEPES, pH 7.4, and 20 μM fenvalerate (DuPont Inc.) to facilitate the analysis of open-channel blockade by Cd2+ (Yamagishi et al., 1997). Well-resolved single-channel opening events were determined using a half-height criterion (Colquhoun and Sigworth, 1985). Amplitude histograms were fitted to the sum of Gaussians using a nonlinear least-squares

172 P Segment Structure of the Na Channel
K channels

| KcsA | turret | pore helix | selectivity filter |
|------|--------|------------|-------------------|
| EERAPGAQLITYPRALWNSVETATGTYGVDLYPVTL |

| hromk |                    |
|-------|--------------------|
| GLTSAFLPSLETQVITGYGFRV |

| hgirk |                    |
|-------|--------------------|
| GFSVSLPSIETETIETYGFRV |

| hKV1.1 |                    |
|-------|--------------------|
| STPDAFWAVYSTMTGVDYMT |

| dShaker |                    |
|--------|--------------------|
| SIPDAFWAVYSTMTGVDYMT |

method. The voltage dependence of Cd$^{2+}$ blockade was used to estimate the fractional electrical distance to the substituted cysteine residue as described previously (Yamagishi et al., 1997). The average current during long openings in the presence of Cd$^{2+}$ was used to quantify the blocked unitary current amplitude.

Pooled data are presented as means ± SD with at least three determinations of the IC$_{50}$ for block by Cd$^{2+}$ and TTX with the exceptions noted as follows. The wild-type IC$_{50}$ for Cd$^{2+}$ block was 2.5 ± 0.2 mM. The mutant channels S387C, W388C, L393C, F742C, H743C, S744C, F749C, G1224C, G1516C, and C1521A exhibited no difference in the IC$_{50}$ compared with the wild-type channel in a single determination that was confirmed by the absence of 50% block of the peak current by 0.5 mM Cd$^{2+}$ in at least two additional cells. The wild-type IC$_{50}$ for TTX block was 17 ± 5 nM. The mutant channels S387C, W388C, A389C, E390C, L391C, F742C, H743C, S744C, S1229C, L1230C, Q1232C, G1516C, N1517C, S1518C, C1521A, F1523C, E1524C, and H1529C exhibited no difference in the IC$_{50}$ compared with the wild-type channel in a single determination that was confirmed by the greater 50% block of the peak current by 50 nM Cd$^{2+}$ in at least two additional cells. Statistical comparisons were made using an analysis of variance (ANOVA) with $P < 0.05$ considered to be significant.

**Molecular Modeling**

Each homologous domain of the Na channel was evaluated by the program PHDhtm (http://www.embl-heidelberg.de/predict-protein), which is a neural network system that predicts the locations of transmembrane helices in integral membrane proteins (Rost et al., 1995). This method uses evolutionary information as input to the network system and detects transmembrane helices with an accuracy of 95%. PHDhtm predictions were first validated with the known structure of the KcsA channel with its default parameters (Hsu et al., 1999): three helical segments were assigned by the program, and they corresponded to TM1, the pore helix, and TM2 of the KcsA. Several inward-rectifying (hROMK1 or hKir1.1, hGIRK1 or hKir3.1) and voltage-gated K channels (hKir1.1, dShaker) were also tested with PHDhtm: the program assigned a helical segment adjacent to the NH$_2$-terminal of the selectivity sequences in each sequence. The sodium channels µ1 and hH1 were then evaluated by input of the entire sequence, and the sequence of each homologous domain using the default setting of PHDhtm. Molecular models were generated using the Insight/Discover suite of programs (Molecular Simulation Inc.). The steepest descent and conjugate gradients were used for energy minimization.

**RESULTS**

**Modeling Predicts Pore Helices in the Na Channel**

We tested the SS1 region of the Na channel P segments for α helical content using PHDhtm, a neural network based program that detects transmembrane helices with 95% accuracy (Rost et al., 1995). We first validated the predictive power of the PHDhtm by testing the program on several proteins of known structure including KcsA (Doyle et al., 1998). The program accurately predicted the transmembrane and pore helices of KcsA (Doyle et al., 1998). PHDhtm predicts α helices involving the italicized residues in the KcsA pore and the P segments of each Na channel domain (Fig. 1). Based on the crystal structure, the second of the two vicinal tryptophans (W68), E71 and T72 in KcsA, are oriented with their side chains accessible to the channel pore (Doyle et al., 1998). PHDhtm predicted helices in sequence-aligned regions of the inwardly rectifying channels hROMK1 (Kir1.1) and hGIRK1 (Kir3.1), the volt-
age-dependent K channels hKv1.1 and Drosophila Shaker (Fig. 1), and the α1C subunit of the Ca channel (unpublished data). Several motifs in the putative helical regions of the P segments of the Na channel are conserved from the KcsA pore helix: a threonine residue one to two positions NH$_2$-terminal to the selectivity filter (except domain II), and conserved aromatic and charged residues in the pore helix. We sought to experimentally test the hypothesis that the regions proximal to the selectivity filter in the Na channel are helical.

Accessibility Pattern to Cd$^{2+}$, MTS Reagents, and TTX

Fig. 1 shows the positions in the Na channel pore studied here. The rat α1 channel P segment sequences are aligned with the bacterial inward-rectifier channel (KcsA). The residues that form the turret and pore helix of the KcsA channel are designated (Doyle et al., 1998). The amino acids underlined with a jagged line are known to be accessible from the external mouth of the channel by thiol-specific modifiers (Chiamvimonvat et al., 1996; Tsushima et al., 1997b; Yamagishi et al., 1997). 3 of the 40 mutations (G1226C, L1231C, and L1522C) did not yield functional channels. To evaluate the possibility that an internal disulfide bond had formed spontaneously to cripple these mutant channels (Benitah et al., 1997), cells were exposed to 1 mM DTT. The application of DTT did not reveal any current (unpublished data).

The permeation phenotype of each expressing mutant channel was characterized by determination of the IC$_{50}$ for block by Cd$^{2+}$ and TTX, accessibility to modification by MTS reagents, and peak current size with cationic substitutions. Fig. 2 shows the sensitivity to externally applied Cd$^{2+}$ of wild-type and mutant Na channels. Eight mutants had enhanced sensitivity to block by Cd$^{2+}$ compared with the wild-type channel: F390C and R395C in domain I; R750C, L746C, and F745C in domain II; Y1227C, L1228C, and I1519C in domain III; and G1526C in domain IV. The lack of deviation from wild-type was confirmed in several other experiments using single doses of Cd$^{2+}$ (0.5 mM) and TTX (50 nM).

**Figure 2.** The effect of external Cd$^{2+}$ and TTX on the cysteine substitution mutants on the NH$_2$-terminal side of putative selectivity filter in α1. (A) Bar plot of the IC$_{50}$ for Cd$^{2+}$ block of each of the mutants that express current. The mutants R395C, R750C, L746C, F745C, Y1227C, L1228C, and I1519C have IC$_{50}$s (mean ± SD) 999 ± 64, 201 ± 10, 815 ± 10, 123 ± 9, 650 ± 106, 292 ± 94, and 173 ± 19 μM (P < 0.05 versus WT, 1,955 ± 207 μM), respectively. (B) Bar plot of the IC$_{50}$ for TTX of each of the mutants that express current. The mutants R395C, F745C, R750C, and L1228C have IC$_{50}$s for TTX block of (mean ± SD) 112 ± 32, 1,600 ± 425, 4,431 ± 661, and 441 ± 288 nM (P < 0.05 versus WT, 13 ± 4 nM), respectively. In cases without standard error bars, full dose–response curves were obtained in individual cells, and the lack of deviation from wild-type was confirmed in several other experiments using single doses of Cd$^{2+}$ (0.5 mM) and TTX (50 nM).
main II; Y1227C and L1228C in domain III; and I1519C in domain IV (Fig. 2 A). It is notable that the mutated residues in the same P segments that exhibit enhanced sensitivity to Cd$^{2+}$/H$_{\text{11001}}$ are either adjacent to one another or separated by three to four amino acids. The easy reversibility of Cd$^{2+}$/H$_{\text{11001}}$ block and the relatively low, micromolar affinity of sensitive channels suggests that single cysteine residues participate in the divalent cation block.

Mutations in the pore of the Na channel also alter sensitivity to block by guanidinium toxins. The toxin molecule is much larger than Cd$^{2+}$/H$_{\text{11001}}$, with more points of contact with amino acids in the channel pore. The IC$_{50}$ for TTX of the wild-type/H$_{\text{9262}}$ channel expressed in tsA201 cells is 17$\pm$0.5 nM, comparable to that in the oocyte expression system (Backx et al., 1992; Satin et al., 1992). Of the eight mutants with altered Cd$^{2+}$ sensitivity, four significantly destabilized TTX block (R395C, F745C, R750C, and L1228C). No mutant channels with unchanged sensitivity to Cd$^{2+}$ block exhibited altered TTX sensitivity (Fig. 2 B). The IC$_{50}$s (mean $\pm$ SD) for TTX block of R395C, R750C, F745C, and L1228C were 112 $\pm$ 32, 4,431 $\pm$ 661, 1,660 $\pm$ 425, and 441 $\pm$ 270 nM, respectively.

MTS reagents selectively modify free cysteinyl side chains in an aqueous environment. External application of saturating concentrations of MTS reagents produced no significant effect on the current through any of the mutant channels; data from R395C are summarized in Fig. 3. To test the MTS accessibility from the inside of the channel, MTSEA or MTSET was applied to the cytoplasmic face of F394C, R395C, and L396C (domain I) and I751C and R750C (domain II). Internal MTS reagents had no effect on the mutant channels (unpublished data).

The lack of effect of MTS reagents may result from an inability of the reagents to form a mixed thiol. Alternatively, the cysteinyl may be modified, but the adduct does not block the permeation pathway and, therefore, does not affect current amplitude. To distinguish between these possibilities, we measured sensitivity to Cd$^{2+}$ block before and after application of MTS reagents. Fig. 3 shows representative experiments for R395C. Application of 1 mM Cd$^{2+}$ blocks $\sim$60% of the current; the subsequent administration of MTSEA, MTSET, and MTSES does not alter the current amplitude nor does it modify the sensitivity to subsequent applications of Cd$^{2+}$ (Benitah et al., 1996, 1997). Similar results were observed with the other Cd$^{2+}$-sensitive mutants (unpublished data).

**Single-channel Analysis**

The voltage dependence of blockade of the unitary current by external Cd$^{2+}$ was used to determine the fractional electrical distance of divalent cation binding. The fractional electrical distance for Cd$^{2+}$ block of cysteine mutants provides the relative location of the substituted thiol in the channel pore. Fig. 4 (top) shows unitary Na$^+$ currents for R395C and R750C, Cd$^{2+}$-sensitive mutants with robust whole-cell expression, in the presence and absence of Cd$^{2+}$. In the absence of Cd$^{2+}$, the single-channel conductances of R395C and R750C differed from the wild-type (50 $\pm$ 0.3 pS in 140 mM [Na$^+$]$_{\text{o}}$) by $<30\%$. In the presence of Cd$^{2+}$, block is rapid and, therefore, appears as a reduction in the single-channel current amplitude. The voltage dependence of block is evident from the change in single-channel current through the mutant channels in the presence and absence of Cd$^{2+}$ over a wide voltage range (Fig. 4, middle). In each case, the reduction in unitary current is more prominent at negative voltages. By plot-
ting the ratio of the blocked and unblocked unitary currents and assuming a single-site model for Cd\textsuperscript{2+} binding (Yamagishi et al., 1997), the fraction of the voltage field traversed by Cd\textsuperscript{2+} to reach its binding site was estimated to be 0.12 and 0.23 for R395C and R750C, respectively (Fig. 4, bottom).

**Ion Selectivity of the Mutant Na Channels**

Several residues in both the third (K1237) and fourth domain (G1530, W1531, and D1532) P segments of the Na channel influence ionic selectivity (Heinemann et al., 1992; Chiamvimonvat et al., 1996; Favre et al., 1996; Perez-Garcia et al., 1997; Tsushima et al., 1997b). We examined all mutants for changes in ion selectivity; only R395C and R750C altered the selectivity of the channel, and R395C altered selectivity most dramatically. We compared the whole-cell conductances in external solutions of different monovalent and divalent cation composition to the Na\textsuperscript{+}/H\textsubscript{+}/K\textsuperscript{+}/Ca\textsuperscript{2+} conductance of the mutants. Fig. 5 (A and B) shows representative R395C currents and current-voltage relationships. Fig. 5 C summarizes the ion-specific conductances compared with Na\textsuperscript{+}. For comparison, the previously reported domain III DEKA mutant, K1237C, is shown (Chiamvimonvat et al., 1996; Perez-Garcia et al., 1997). R395C is more permeable to NH\textsubscript{4}\textsuperscript{+}, K\textsuperscript{+}, and Ca\textsuperscript{2+} than the wild-type channel. R750C exhibits enhanced permeability to NH\textsubscript{4}\textsuperscript{+} and modest K\textsuperscript{+} permeability, but does not support divalent cation flux. The selectivity phenotype of F745C is similar to the wild-type.

**DISCUSSION**

The P segments contain the major determinants of permeation in voltage-gated ion channels. In the Na channel, the DEKA ring is a central, but not the only element that mediates conductance and selectivity (Chiamvimonvat et al., 1996; Tsushima et al., 1997b). Amino acid residues on both the NH\textsubscript{2}- and COOH-terminal sides of the DEKA ring are accessible from the extracellular side of the channel pore, but no P segment residues have been demonstrated to contribute to the inner mouth of the Na channel (Chiamvimonvat et al., 1996; Perez-Garcia et al., 1996; Tsushima et al., 1997b; Yamagishi et al., 1997). We have previously shown that cysteine mutants immediately NH\textsubscript{2}-terminal of the DEKA ring in domains III (F1236C and T1235C) and IV (S1528C) are accessible from the outer pore mouth. The data suggest that these P segments descend then ascend in the outer pore but, unlike the K channel, do not span the selectivity region (Yamagishi et al., 1997).

![Figure 4. Single-channel Cd\textsuperscript{2+} block of R395C and R750C.](image-url)
ture referred to as the pore helix. Side chain accessibility to the aqueous phase of SSI residues was gauged by a combination of cysteine substitution mutagenesis, and sensitivity to block or modification by hydrophilic, thiol-specific modifying reagents. The probes used provide complementary structural information. Alterations in external Cd\textsuperscript{2+} block sensitivity and MTS modification of the current imply side chain accessibility in the outer mouth of the pore. Modest changes in TTX block may be associated with global changes in pore structure with retained specific channel–toxin interaction sites. Dramatic changes in toxin affinity are more likely to result if a specific channel–toxin interaction is disrupted (Hidalgo and MacKinnon, 1995). In this regard, it is interesting to note that only mutations in domain II in this region of the P segment produce dramatic (more than three orders of magnitude) changes in sensitivity to TTX block (Fig. 2B).

In each domain, at least one substituted thiol side chain is accessible to the extracellular pore. In domain I, the side chains of both F390C and R395C (separated by four residues) are accessible. In domain II, F745C, L746C, and R750C exhibit side chain accessibility. In domain III, Y1227C and L1228C have enhanced sensitivity to Cd\textsuperscript{2+} block, whereas G1226C and L1231C have an indeterminate phenotype since they do not express functional channels. Similarly, in domain IV, the mutant I1519C exhibits side chain accessibility to Cd\textsuperscript{2+} and is three residues NH\textsubscript{2}-terminal from the nonfunctional mutant L1522C. In the context of previous mutagene-

\textbf{Figure 5.} Selectivity of the cysteine mutants with altered Cd\textsuperscript{2+} and TTX affinities. Peak inward currents (A) and current-voltage relationship (B) through R395C in the presence of different cations. (C) Plot of the ratio of the whole-cell conductance in 140 mM Li\textsuperscript{+}, 140 mM NH\textsubscript{4}\textsuperscript{+}, 140 mM K\textsuperscript{+}, and 70 mM Ca\textsuperscript{2+} \((G_x)\) compared with that in 140 mM Na\textsuperscript{+} \((G_{Na})\). The selectivity of the wild-type channel and the selectivity filter mutant K1237C are shown for comparison.
Figure 6. Structure of Na channel SSI region of the P segments. (top) Helical wheel representations of the residues mutated in this study. The residues in bold italics are accessible from the aqueous pore; those enclosed in a box do not express current and thus have an indeterminate phenotype. The cysteine mutants that alter Ca$^{2+}$/TTX block sensitivity or have an indeterminate phenotype are on the same side of the helix. (bottom) Molecular model of the putative pore helices in the Na channel pore, views from the top (left) and side (right) are shown. The domains are positioned to approximate the distances between selectivity filter residues. The pore helices are rendered in ribbon format, the DEKA ring in yellow stick format, and accessible (green) or nonfunctional (salmon) positions in ball-and-stick format.
sensitivity data in adjacent regions of the P segments (Yamagishi et al., 1997), side chains that are accessible or indeterminate are observed every three to five positions NH2-terminal to the DEKA ring in domains II–IV (Figs. 1 and 6). This periodicity of side chain accessibility in the SS1 regions is consistent with \( \alpha \) helices.

Helical wheel representations of the SS1 regions demonstrate that the accessible or indeterminate side chains lie on the same face of the predicted \( \alpha \) helices with the exception of G1226 (Fig. 6). The secondary structure of amino acid residues on the NH2-terminal side of DEKA are distinct from the random coil structure predicted by mutagenesis of amino acids on the COOH-terminal side of the ring (Chiamvimonvat et al., 1996; Perez-Garcia et al., 1996; Tsushima et al., 1997b).

The detailed pattern of accessibility differs for each of the residues that alter channel interaction with the structural probes. The mutations R395C (domain I), F745C, R750C (domain II), and L1228C (domain III) increase the affinity for Cd\(^{2+}\) block and are resistant to TTX. However, external application of MTS reagents does not alter the current through these mutant channels. R395C and R750C, but not F745C or L1228C, alter cation selectivity of the channel (Fig. 5). R395 and R750 are at homologous positions by sequence alignment in domains I and II, respectively. It is possible that they form a charge ring in the external pore that alters permeation. However, R395C and R750C do not effect permeation by a simple electrostatic mechanism. Neutralization of positive charges in the external pore would be expected to increase channel conductance; neither mutant increases conductance compared with wild type, in fact, R750C is \( \sim 30\% \) smaller than R395C or wild type (Fig. 4). A simple electrostatic mechanism cannot explain the change in ion selectivity observed with these two mutations (Fig. 5). Furthermore, an electrostatic impediment to TTX binding is inconsistent with the data: a reduction in the net positive charge of the pore, all else being equal, should increase TTX sensitivity. Instead, these two charge-reducing mutations decrease the channel sensitivity to block by TTX, perhaps by changing the interaction of other toxin channel contact sites (Noda et al., 1989; Terlau et al., 1991; Lipkind and Fozzard, 1994; Chiamvimonvat et al., 1996; Perez-Garcia et al., 1996; Chen et al., 1997).

It is possible that these residues are involved in an electrostatically mediated stabilization of the pore-forming salt bridges with the acidic filter residues in domains I (D400) and II (E755). Of course more global changes in the pore structure cannot be ruled out with certainty, but the relative preservation of Na\(^{+}\) selectivity (Fig. 5 C) argues against this possibility.

Another mutation that alters both Cd\(^{2+}\) and TTX sensitivity is replacement of aromatic phenylalanine in domain II (F745C). Both mutations in domain II that alter Cd\(^{2+}\) sensitivity (F745C and R750C) decrease TTX affinity by nearly two orders of magnitude. If F745 and R750 participate in the formation of a pore helix, the cysteine substitutions at these positions disrupt an important interaction between the channel and toxin. F745C produces no change in channel selectivity and is not modified by MTS reagents. Similarly, L1228C increases affinity for Cd\(^{2+}\) block and makes the channel resistant to TTX block. The magnitude of the destabilization of TTX block is similar to that exhibited by the domain I mutant R395C and less than that of the domain II mutants F745C and R750C. The remainder of the cysteine mutations, Y1227C (domain III) and I1519C (domain IV) increase Cd\(^{2+}\) affinity without affecting toxin block or MTS binding.

The IC\(_{50}\) for Cd\(^{2+}\) block of all of the mutants is not affected by prior addition of MTS reagents. This suggests that the cysteine mutants rather than being modified but not altering current are, instead, not modified by MTS reagents. These data imply that some parts of the P segments in the external pore are not accessible to modification by external MTS reagents, which require a reaction sphere of \( \sim 6 \) A.

The pattern of side chain accessibility in the pore helix region has been studied in a number of K channels. As in the case of the Na channel, and as predicted by the KcsA structure, hydrophobic residues at the NH2-terminal side of the pore helix appear to have their side chains exposed using Ag\(^{+}\) (Lu and Miller, 1995) or charybotoxin (Ranganathan et al., 1996) as structural probes. In contrast, there is less experimental data to assign the location of the side chains of the residues at the COOH-terminal side of the helix. However, cysteine scanning mutagenesis followed by MTS modification generally has shown that the residues in this region of the K channel pore are inaccessible (Kurz et al., 1995; Pascual et al., 1995). Residues at or near the COOH-terminal side of the putative pore helix are involved in the determination of sensitivity to block by internal TEA (Yellen et al., 1991). The absence of an internally accessible site flanking the pore helices of the Na channel suggests a different orientation compared with K channel pore helices.

**Summary**

Our data demonstrate that the P segments on the NH2-terminal side of the DEKA residues of the Na channel participate in formation of the outer pore. The P segments do not span the selectivity filter; none of the cysteine substitutions are accessible from the internal mouth of the channel. The pattern of side chain accessibility is consistent with pore helices similar to that seen in the KcsA channel pore crystal structure and side chain accessibility in other K channels. Two of the mutations in domain II (F745C and R750C) that alter
Cd\(^{2+}\) block dramatically reduce TTX sensitivity, and may be sites of direct toxin–channel interaction. The data are consistent with previous studies that demonstrate the importance of domains I and II in guanidinium toxin binding and block (Noda et al., 1989; Terlau et al., 1991; Backx et al., 1992; Satin et al., 1992; Lipkind and Fozzard, 1994; Chiamvimonvat et al., 1996; Perez-García et al., 1996; Chen et al., 1997). Arginine residues in the first and second domains (R395 and R750) alter cation selectivity, and in conjunction with the DEKA ring and residues in domain IV, comprise the selectivity determinants of the Na channel.

Our data and previous work (Chiamvimonvat et al., 1996; Perez-García et al., 1996; Chen et al., 1997; Tsushima et al., 1997a,b; Yamagishi et al., 1997) provide experimental evidence to support the hypothesis that the SSI regions of the Na channel form an α helix-turn-coil motif (Fig. 6, bottom). The precise location of the end of the helix and beginning of the turn are not defined; however, modeling predicts that 8–10 amino acids preceding the selectivity filter residue in each domain, have helical character. Overall, this region is hydrophobic, and helical packing is likely to be thermodynamically favorable for this water-accessible part of the protein. The preference for acidic residues to flank the NH\(_2\) terminus and basic residues the COOH terminus (Lipkind and Fozzard, 2000) suggests the possibility that the NH\(_2\) termini of the α helices may extend beyond the region studied; however, in domains III and IV, this would mean the incorporation of one or more glycines into the helices.

The KcsA channel appears to be an evolutionary forerunner of the voltage-dependent family of K channels (MacKinnon et al., 1998). In turn, Na and Ca channels are likely to have evolved from voltage-dependent K channels by gene duplication (Strong et al., 1993). Such evolutionary relationships may explain the remarkable homologies of the mammalian Na channel pore with that of a bacterial K channel. It is likely that the pore helix is essential for the integrity of the transmembrane pore common to all ion channels (Doyle et al., 1998). However, it is the substantial differences in structural detail that mediate the unique functions of the Na channel pore.

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