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Abstract

Although the interiors of membrane and water-soluble proteins are similar in their physicochemical properties, membrane proteins differ in having larger fractions of hydrophobic residues on their exteriors. Thus, it should be possible to water-solubilize membrane proteins by mutating their lipid-contacting side chains to more polar groups. Here, a computational approach was used to generate water-soluble variants of the potassium channel KcsA. As a probe of the correctness of the fold, the proteins contain an agitoxin2 binding site from a mammalian homologue of the channel. The resulting proteins express in high yield in*Escherichia coli* and share the intended functional and structural properties with KcsA, including secondary structure, tetrameric quaternary structure, and tight specific binding to both agitoxin2 and a small molecule channel blocker.

Disciplines

Biochemistry | Organic Chemistry | Physical Chemistry

Computational design of water-soluble analogues of the potassium channel KcsA

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Although the interiors of membrane and water-soluble proteins are similar in their physicochemical properties, membrane proteins differ in having larger fractions of hydrophobic residues on their exteriors. Thus, it should be possible to water-solubilize membrane proteins by mutating their lipid-contacting side chains to more polar groups. Here, a computational approach was used to generate water-soluble variants of the potassium channel KcsA. As a probe of the correctness of the fold, the proteins contain an agitoxin2 binding site from a mammalian homologue of the channel. The resulting proteins express in high yield in *Escherichia coli* and share the intended functional and structural properties with KcsA, including secondary structure, tetrameric quaternary structure, and tight specific binding to both agitoxin2 and a small molecule channel blocker.

he 3D structures of tens of thousands of water-soluble proteins have been solved, but the structures of membranesoluble proteins have proven much more difficult to determine. The problems associated with membrane proteins include their limited levels of expression, low stability in detergent-solubilized forms, and greater difficulty in obtaining high-resolution diffraction quality crystals. Thus, we and others have sought to develop a broadly applicable automated strategy for the preparation of water-soluble mutants of membrane proteins, which could be obtained in larger quantities for a variety of high-resolution biophysical and drug discovery studies. Initial work yielded good results for a simple transmembrane coiled-coil (1-3), but attempts to design water-soluble versions of larger proteins such as bacteriorhodopsin led to misfolded proteins with very limited solubility in water (4). Here, we demonstrate the success of a previously undescribed computational approach for the design of a water-solubilized version of a bacterial ion channel with a transplanted mammalian toxin-binding site. These findings have fundamental implications concerning the stabilization of membrane vs. water-soluble proteins (5) as well as practical implications for the design of water-soluble analogues of a variety of biologically interesting membrane proteins.

The structures of many membrane proteins are beginning to appear, allowing one to infer features common to entire families. In particular, the structures of bacterial ion channels (6-12) have provided insight into their mammalian counterparts, and the structure of rhodopsin (13) has served as a prototype for the entire family of seven-transmembrane G protein-coupled receptors. It would be advantageous to use what structural information is available for a given membrane protein to obtain water-soluble versions that retain their structure, oligomerization state, and essential ligand-binding properties. As a target for such an effort, we have chosen the bacterial KcsA potassium channel because of its available structure (6, 12), its biochemical characterization, and the interest in this family of channel proteins. Furthermore, the external vestibule of the ion-conducting pore of KcsA has been mutated to the corresponding residues in a mammalian channel (Q58A, T61S, and R64D) to allow binding of agitoxin2 (AgTx₂), resulting in a protein (designated here as tKcsA) that binds AgTx₂ (14). This system extends multiple clearly defined criteria for the successful design of a water-soluble version of this protein, which should: (*i*) be expressed at high level in a water-soluble form; (*ii*) show the correct helical secondary structure; (*iii*) associate to form tetramers; (*iv*) bind AgTx₂ with high affinity, specificity, and in the appropriate stoichiometry; and (*v*) bind small molecule channel blockers such as tetraethylammonium chloride (TEA). The designed water-soluble variants of tKcsA are referred to as WSK-1, -2, and -3.

Methods

Protein Design. A statistical entropy-based formalism has been developed for identifying amino acid probabilities from a given backbone structure (15, 16). This method takes as input a target structure, in this case a high-resolution structure of KcsA (Protein Data Bank ID code 1k4c), and energy functions that quantify sequence-structure compatibility. The output is the set of site-specific probabilities of the amino acids compatible with the structure. The site-specific probabilities of the amino acids and their discrete side chain conformational states (rotamers) are determined by maximizing an effective entropy function subject to simultaneous constraints on both the overall energy as determined by an atom-based potential and the value of an effective solvation score ("environmental energy"). In the calculation, all 20 amino acids and up to 10 of their side chain conformations [rotamer states (17)] were considered at each site where mutations were permitted. This yields a total of 129 identity-rotamer states for each variable position. C4 subunit symmetry was imposed, reducing the complexity (total number of possible sequence-rotamer combinations) from 129140 to 12935 for the first of the recursive calculations discussed below (18).

To quantify hydrophobicity and solvation effects, an environmental potential is used (16), where the local C β density about each residue is used to quantify its degree of exposure to solvent. The environmental energy for the entire protein (E_{env}) and for the buried residue sites ($E_{env,b}$) are constrained to the average values of water-soluble proteins in the calculation (-46.0 and -24.6). The identities of buried sites are constrained in these calculations, and the decrease in E_{env} (from +20 to -46) largely results from the mutation of exposed hydrophobic residues to more hydrophilic amino acids.

Nonbonded interactions involving the side chains are calculated by using the AMBER 3A force field (19) with a modified hydrogen-bonding term (20) and a distance-dependent dielectric constant ($\varepsilon = 4$ r). To address unfolded states, a reference

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Abbreviations: AgTx₂, agitoxin2; TEA, tetraethylammonium chloride; AgTx₂-DNP, AgTx₂ with 2,4-dinitrophenyl-Ala; E_{env} , environmental energy.

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energy $\gamma_{\text{ref}}(\alpha)$ for each amino acid is introduced into the energy E_c to represent the effects of the denatured state. The energy is calculated as a "free energy" of each amino acid in its *N*-acetyl-N'-methylamide derivative with averaging over multiple backbone and rotamer states. This averaging involves a sum over possible rotamers and possible backbone configurations, approximated by varying each of the backbone ϕ and φ angles in increments of 10°. This crudely approximates an average over extended unfolded states. The reference energies of each amino acid may then be estimated by using

$$\gamma_{\rm ref}(\alpha, \beta_{\rm ref}) = -\beta_{\rm ref}^{-1} \ln(z_{\rm ref}(\alpha, \beta_{\rm ref})/z_{\rm ref}(G, \beta_{\rm ref}))$$
$$z_{\rm ref}(\alpha, \beta_{\rm ref}) = \sum_{\phi, \phi, k} \exp(-\beta_{\rm ref} \varepsilon_{\rm ref}(\phi, \phi, r_k(\alpha))),$$

where γ_{ref} is the conformational energy in a particular conformation of the *N*-acetyl-*N'*-methylamide derivative of the amino acid, as determined by using the molecular potential. Here $\beta_{ref} = 1/k_{\rm B}T$, where $k_{\rm B}$ is Boltzmann's constant and T is a temperature appropriate for the conformation sampling of side chain and backbone conformations (e.g., T = 300 K). Reference energies are expressed relative to Gly (16). In the statistical formalism of *Scads* (statistical computationally assisted design strategy), accompanying the interatomic potential energy is a corresponding effective temperature $1/\beta$. The probabilities used in sequence identification were determined for $\beta = 0.5$ mol/kcal; at this value, we find the sequence properties are robust with respect to slight variation in backbone structure (16).

WSK Expression and Purification. The gene encoding WSK-1 was synthesized with *Pfu* polymerase by using three fragments. The final WSK-1 fragment was cloned into the pET-24a(+) vector (Novagen) at the NdeI and XhoI restriction sites, expressing no tag. Mutants WSK-2 and -3 were generated by using QuikChange (Stratagene). The WSK proteins were expressed in BL21(DE3) cells (Novagen) in LB for 4 h after induction with 1 mM isopropyl- β -D-thiogalactoside. Cells were harvested by centrifugation at 4°C and 3,450 $\times g$. Cell pellets were lysed by French press at 1,500 psi in lysis buffer containing 10 mM Tris[hydroxymethyl]aminomethane hydrochloride, pH 7.0, and 1 mM EDTA. Cell extracts were loaded onto a 40-ml Q Sepharose (Amersham Pharmacia Biosciences) column in the lysis buffer and eluted with a step gradient of the lysis buffer containing 0–500 mM KCl. Fractions containing WSK were pooled, concentrated to 50 ml, and further purified by reverse-phase HPLC on a Vydac (Hesperia, CA) C4 preparative column by using a linear gradient of water and acetonitrile containing 0.1% trifluoroacetic acid. Purity was verified by analytical reverse-phase HPLC, matrix-assisted laser desorption ionization time-of-flight mass spectrometry, and 4-12% Bis-Tris SDS/PAGE gels. The molar extinction coefficient ($\varepsilon_{280nm} = 17,781 \text{ M}^{-1} \cdot \text{cm}^{-1}$) was determined by the difference in absorbance of a sample in 0 and 6 M guanidine HCl, by using the calculated extinction coefficient at 280 nm as a starting point.

AgTx₂ Synthesis and Purification. AgTx₂ (4,097 Da) was chemically synthesized as a C-terminal carboxyamide on a 0.25 mmol scale by using an Applied Biosystems model 433A solid phase peptide synthesizer (Perkin–Elmer) with standard fluorenyl-methoxycarbonyl amino acid chemistry. The tripeptide gly-gly-*N*-2,4-dinitrophenyl-Ala chromophore was coupled to the N terminus of half of the resin. AgTx₂ with 2,4-dinitrophenyl-Ala (AgTx₂-DNP) has a calculated $\varepsilon_{356nm} = 11,343 \text{ M}^{-1} \cdot \text{cm}^{-1}$, and $\varepsilon_{280nm} = 4,520 \text{ M}^{-1} \cdot \text{cm}^{-1}$. Peptides were washed on the resin with dimethylformamide and ether, cleaved for 3.5 h with

trifluoroacetic acid/water/ethanedithiol/triisopropylsilane (94.5/2.5/2.5/1) vol/vol/vol, and precipitated and washed with cold ether. Purification proceeded by reverse-phase HPLC by using a preparative C4 column (Vydac) and a linear gradient of the appropriate buffers. AgTx2 required oxidative refolding to attain the proper folded state after HPLC. A series of redox buffers containing increasing proportions of oxidized (GSSG) and reduced glutathione (GSH) were made. The concentration of GSSG was determined by UV/Vis absorbance by using ϵ_{282nm} = 273 $M^{-1} \cdot cm^{-1},$ whereas GSH was monitored by titration with Elmans reagent, following absorbance at 412 nm. Optimal conditions were identical for both labeled and unlabeled peptides, and the final reaction components were determined to 125 μ M toxin in the presence of air and 100% GSH (6 mM). The progress of the reaction was followed by using analytical reverse-phase HPLC and comparison of the magnitude of the properly folded peak to the misfolded peaks. Properly folded toxins were repurified by reverse-phase HPLC and tested for function.

Solution Measurements. CD spectra were collected on an Aviv 62DS (Aviv Associates, Lakewood, NJ) spectropolarimeter, by using a 1-mm path-length quartz cuvette, with protein at 44 μ M, in 20 mM potassium phosphate, pH 7.0/100 mM KCl/1 mM EDTA. For ultracentrifugation, samples were centrifuged at 15,000 rpm in 20 mM potassium phosphate, pH 7.0/100 mM KCl/1 mM EDTA by using a Beckman XL-I (Beckman Coulter) analytical ultracentrifuge. Equilibrium dialysis to determine the stoichiometry of toxin binding to WSK-3 was performed by using 0.5 ml of centrifuge tubes and a dialysis membrane with a 10-kDa molecular weight cutoff. After reaching equilibrium, the concentrations of AgTx₂-DNP and WSK-3 were determined by absorbance at 356 and 280 nm, respectively.

The extent of binding of AgTx₂-DNP (10 μ M) to WSK-3 (10 μ M, tetramer) was assessed at various concentrations of TEA and tetramethylammonium ion by ultracentrifugation at $16,385 \times g$. Because we expected some salt dependence to the binding interaction, we also measured the extent of binding in the presence of the same concentrations of KCl. The fraction of free AgTx₂-DNP was determined from the absorbance at 360 nm at the top of the cell (at this speed, the WSK-3 is essentially fully depleted, but the AgTx₂-DNP has not significantly sedimented). The extent of binding, relative to the KCl control, was plotted and the IC₅₀ determined (Fig. 5). The approximate K_{diss} for binding to TEA is determined from the relationship $IC_{50} = K_{TEA} [f_{AgTx2-DNP}]/K_{AgTx2-DNP}$, where f_{AgTx2-DNP} is defined as the free AgTx2-DNP concentration, and K_{TEA} and $K_{\text{AgTx2-DNP}}$ are the dissociation constants for WSK-3 with TEA and AgTx₂-DNP respectively. This relationship holds approximately for $[f_{AgTx2-DNP}] \gg K_{AgTx2-DNP}$, where $[f_{AgTx2-DNP}] = 5 \ \mu M$ and $K_{AgTx2-DNP} \le 2 \ \mu M$ (because tight binding is observed in the equilibrium dialysis with 10 μ M total toxin concentration).

Results and Discussion

Design of WSK. The side chains of the transmembrane helices of tKcsA were stripped to the backbone and then rebuilt in accordance with the requirements for function, stability, and water solubility. We retained the pore-lining residues, the extracellular loops, the intracellular region, and the residues required for binding to $AgTx_2$. Furthermore, we retained the buried residues, leaving 35 membrane-exposed side chains [>40% accessible to a probe with a radius of 1.4 Å (21)] per protomer as targets for design. These positions were chosen by using the computational design algorithm scads.

A key input into these calculations is an environmental energy, a database-derived quantification of solvation and

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Fig. 1. E_{env} (16) vs. chain length. E_{env} quantifies solvation and hydrophobic effects and has been parameterized by using a database of 500 soluble proteins (small circles) (16). For each protein structure, $E_{env} = \sum_i e(a_i, r_{a,i}, \rho_{C\beta})$ where for the *i*th residue a_i , $r_{a,i}$, and $\rho_{C\beta}$ are the amino acid, side chain conformation, and local density of β carbons. Also shown is the E_{env} for WT KcsA (open circle) and the value E_{env} that was used as a constraint in the sequence calculations (black circle).

hydrophobic effects (16). The value of this environmental score for WT KcsA is +20, which is well outside the range observed for soluble proteins of this size, due to the large number of exposed hydrophobic residues. Thus in calculations to determine water-soluble sequences, the environmental energy was constrained to the value expected for soluble proteins of this size, -46 (Fig. 1).

Scads generates profiles describing the site-dependent probabilities of the amino acids at each position allowed to mutate. A unique sequence was selected from the computed probabilities by using recursive calculations. The set of 35 exposed residues was initially targeted for variation. The identities of the remaining residues were fixed at WT, and their side chain conformations were permitted to vary to accommodate mutations (15, 16). A sequence was selected comprising the most probable amino acids at each site with the exception of several sites where alternate amino acids with appreciable probability were selected: W26E and G43A were chosen for their favorable helical propensities (22, 23) and S44, S69, and V106 were kept at WT. The WT amino acid was selected at 8 of the 35 sites. A second calculation was then performed. In the presence of the resulting designed sequence from the first iteration, an additional eight exposed hydrophobic sites were targeted for mutation. The WT was most probable at L36, V39, L40, I60, and L105; hydrophobic residues were probable at F116, and the WT was retained. The mutation V93E was most probable, and the L24D mutation was selected over the more probable W and F due to its greater polarity. At position 104, Gly and Ala had comparable probabilities, and Ala was chosen for its aqueous helix propensity. The resulting sequence was compared to 47 aligned K⁺ channel sequences from eight prokaryotic and eukaryotic organisms (BLAST, National Center for Biotechnology Information database).

The resulting sequence (WSK-1) had 29 mutations relative to tKcsA (Fig. 2). Preliminary studies discussed below indicated that this soluble protein bound toxin but formed highorder oligomers. The model of WSK-1 showed two exposed hydrophobic patches in the redesigned transmembrane helices, which might mediate aggregation. This problem was addressed





Fig. 2. Depiction of KcsA and WSK-3. Only the side chains of outer helices (22–71), inner helices (89–124), and cytoplasmic and extracellular residues are shown. Side chains are colored based on frequency of occurrence in the apolar section of the lipid bilayer beginning with most probable: Ala/Ile/Leu/Val (dark green), Gly/Met/Thr (light green), and Pro/Ser/Trp/Tyr (light blue). Lys/Arg/Gln (dark blue), Asp/Glu (red). (a) KcsA. Solvent-exposed residues of KcsA allowed to vary in the design are depicted along the inner and outer helices (light green, dark green). Also shown are the cytoplasmic and extracellular residues, unchanged in our design. (b) KcsA structure with side chains of mutated residues removed. Extracellular and cytoplasmic residues held constant from KcsA to WSK are rendered. Buried residues within the interior of the structure are not shown. (c) WSK-3. Sequences of KcsA and WSK-3 are also shown, where colored residues were mutated, whereas those in black were not. Mutations to tKcsA are shown in a gray box [made by using PYMOL (DeLano Scientific, San Carlos, CA).

in a new round of computational design varying only these exposed hydrophobic regions, resulting in WSK-2, which contained the additional mutation L116R, and WSK-3, which contained both L81R and L116R.

The WSK sequences result from the simultaneous imposition of effective energetic constraints on the solvation properties and the interatomic interactions of the flexible amino acid side chains. One might expect that simply mutating exposed hydrophobic residues to polar or charged amino acids might be sufficient to confer solubility, but such approaches often yield misfolded proteins (4). The calculations provide sequences with sterically consistent nontrivial patterning of amino acid identities. Among the polar mutations that were introduced, complementary charge interactions on the protein surface are apparent (see Fig. 2). Although the overall goal is to produce a water-soluble structure, it must also retain sufficient hydrophobic interactions to drive protein folding in an aqueous environment. Thus, some exposed hydrophobes are retained, and one mutation actually yields increased hydrophobic character (S102V). Others have noted the importance of careful consideration of interactions between surface residues in protein design (24).

Secondary and Quaternary Structure of WSK Variants. WSK variants were expressed in high yield (20 mg/ml) and in soluble form in *Escherichia coli*. The CD spectra of the purified proteins were consistent with the expected secondary structure, showing minima at 208 and 222 nm ($[\theta_{222}] = -14,600$ deg·cm²·dmol⁻¹). The computed α -helical content was $\approx 50\%$ (25), in good agreement with the helical content of 60% observed for KcsA (Protein Data Bank code 1k4c).

Size exclusion chromatography was used to determine the aggregation state of WSK variants. WSK-1 eluted as three peaks: one peak consistently eluted as a large aggregate in the void volume, whereas the observed molecular weights of the other two peaks were 10,600 and 50,100 Da (Fig. 3a), in good agreement with the expected masses for the monomer (11,433 Da) and tetramer (45,732 Da), respectively. Preliminary experiments (not shown) showed that both the tetramer and the higher-order aggregate bound AgTx₂ in the proper stoichiometry, suggesting that the void volume peak might consist of loosely associated but otherwise properly folded tetramers. To test this possibility, the molecular weight distribution was measured in the presence of 6 M urea, a concentration lower than that required to unfold the protein (as determined from the loss of secondary structure followed by CD spectroscopy). Under this condition, only the tetrameric and monomeric peaks were observed (Fig. 3b).

WSK-1, -2, and -3 contain progressively fewer apolar side chains in the redesigned transmembrane helices; these mutants also show a progressively smaller fraction of aggregated protein eluting in the void volume. Even in the absence of urea, WSK-3 elutes primarily as a tetramer and shows only a small peak near the position expected for a 12-mer, presumably a trimer of tetramers (Fig. 3c and d). Thus, iterative mutagenesis guided by computation and experiment was able to minimize the nonspecific aggregates of tetramers seen in WSK-1.

AgTx₂ Binding. Size exclusion chromatography was used to demonstrate that WSK-3 specifically bound to AgTx₂. For these experiments, we synthesized a variant of AgTx₂ with AgTx₂-DNP at its N terminus, allowing the toxin to be detected from its absorbance at 356 nm ($\varepsilon_{356nm} = 11,340 \text{ M}^{-1} \cdot \text{cm}^{-1}$). When AgTx₂-DNP was chromatographed through the Superdex 200 column, it eluted as a monomer, but when incubated with WSK variants (10 μ M, assuming a functional tetramer),



Fig. 3. Analytical gel filtration chromatography of WSK-1 (20 μ M), WSK-1 (20 μ M, 6 M Urea), WSK-2 (100 μ M), and WSK-3 (100 μ M) by using a 25-ml Superdex 200 column (Amersham Pharmacia Biosciences) in 20 mM K₂PO₄, pH 7.0/100 mM KCl/1 mM EDTA at 1 ml/min flow rate. The approximate volumes of elution for the monomer (Mon), tetramer (Tet), 12-mer, and high-order aggregate (Agg) are indicated. The column was calibrated by using blue dextran, BSA (66 kDa), ovalbumin (43 kDa), and carbonic anhydrase (29 kDa).

it coeluted with the tetrameric and higher-order aggregate peaks of WSK-1, -2, and -3 (data not shown) in approximately the expected molar ratio of one $AgTx_2$ -DNP for every four molecules of WSK.

Equilibrium analytical ultracentrifugation confirmed the selective association of WSK-3 with AgTx₂-DNP (Fig. 4). WSK-3 sedimented in a concentration-dependent manner, and the traces are well defined by a monomer–tetramer–12-mer equilibrium (Fig. 4*a*), as expected from size exclusion chromatography. Under the same conditions, AgTx₂-DNP shows little sedimentation, consistent with its monomeric molecular weight (4,435 Da). When incubated with WSK-3, AgTx₂-DNP cosediments with the water-solubilized channel (Fig. 4*b*), as determined by the DNP absorbance. At a mol ratio of one AgTx₂-DNP per WSK-3 tetramer (17 μ M), there was essen-



Fig. 4. Equilibrium sedimentation analytical ultracentrifugation of WSK-3 and AgTx₂-DNP. (a) 17 μ M WSK-3 monitored at 280 nm (o). Data were fit to a monomer–tetramer–12-mer equilibrium by using a macro in IGOR-PRO (WaveMetrics, Lake Oswego, OR). (b) AgTx₂-DNP (17 μ M) in the presence (red circle) and absence (green circle) of 17 μ M WSK-3 tetramer, monitored at 360 nm to detect only the AgTx₂-DNP. (c) Control with BSA (36 μ M) plus AgTx₂-DNP (17 μ M) monitored at 280 nm (blue square) to monitor the BSA, and at 360 nm (green square) to monitor AgTx₂-DNP.

tially no free AgTx₂-DNP (26), indicating that the dissociation constant for binding was $\leq 2 \mu M$ (i.e., 10-fold tighter than the total toxin concentration). By contrast, when AgTx₂-DNP was centrifuged in the presence of BSA (36 μ M), no association was observed (Fig. 4*c*), demonstrating the specificity of the AgTx₂-WSK-3 interaction.

The stoichiometry of toxin binding was determined by equilibrium dialysis by using two starting conditions: one with equimolar AgTx₂-DNP and WSK-3 tetramer (10 μ M), as well as one with a 2-fold excess of AgTx₂-DNP. In both experiments, the ratio of AgTx₂-DNP:WSK-3 monomer was 1:4 within experimental error (1:4.4 and 1:4.1, respectively). Control experiments showed no binding of AgTx₂-DNP to either BSA or carbonic anhydrase (Table 1).

TEA Competition. To determine whether small molecule channel blockers also bind specifically to WSK-3, we measured the ability of TEA to compete with AgTx₂-DNP for binding to WSK-3. This small molecule blocker has been shown to compete with charybdotoxin (a homologue of $AgTx_2$) for binding to the extracellular opening of the pore (27). TEA induces dissociation of AgTx2-DNP from WSK-3 in a concentration-dependent manner, whereas the control tetramethylammonium ion [TMA, which does not bind to K^+ channels (28)] showed no significant concentration-dependent competition (Fig. 5). From the concentration of TEA required to induce 50% total toxin dissociation, we calculate a $K_{\text{diss}} \leq 80$ mM for TEA (assuming a $K_{\rm diss} \leq 2 \ \mu M$ for binding of AgTx₂-DNP). This value agrees well with the reported K_{diss} of 3.2–22 mM (29) for the native channel. TEA binds to the outer vestibule of the channel KcsA channel. The finding that TEA binds to WSK-3 strongly suggests that this region of the protein

Table '	1.	Equilibrium	dialysis	of	WSK-3	and	AgTx ₂ -DI	٧P
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	[Tox]	
	bound,	
Experiment	μM	Stoichiometry
WSK-3 _{mon} (40 μ M) + Tox (10 μ M)	9.0	$[WSK-3]/[Tox]_{bound} = 4.4$
WSK-3 _{mon} (40 μ M) + Tox (20 μ M)	9.7	$[WSK-3]/[Tox]_{bound} = 4.1$
BSA (20 μ M) + Tox (44 μ M)	0	
CA (47 μ M) + Tox (23 μ M)	0	

Initially, WSK-3 and AgTx₂-DNP (Tox) are on side *a*, whereas buffer is on side *b*. At equilibrium, the concentration of bound toxin is given by $[Tox]_{bound} = [Tox]_a - [Tox]_a - [Tox]_a - [Tox]_b$.

is properly folded. Although this finding alone does not preclude the possibility of misfolding elsewhere in the protein, the additional CD, size exclusion chromatography, and toxinbinding data together suggest that the protein has a structure similar to the WT channel.

Conclusion

WSK-3 has successfully recapitulated the target properties of tKcsA, including its ability to bind a protein toxin and a small molecule blocker. These findings should be of practical utility for the design of water-soluble analogues of membrane proteins that have proven difficult to obtain in large quantities for biophysical studies or drug screening. The method requires only a medium-resolution structure for computational modeling. Thus, it should be possible to extend this work to other ion channels and G protein-coupled receptor proteins (30) using homology models as starting points.



Fig. 5. The small-molecule blocker TEA competes with $AgTx_2$ -DNP for binding to WSK-3. A competition curve for tetramethylammonium ion is also shown.

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