Structure of Functional *Staphylococcus aureus* α-Hemolysin Channels in Tethered Bilayer Lipid Membranes

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ABSTRACT We demonstrate a method for simultaneous structure and function determination of integral membrane proteins. Electrical impedance spectroscopy shows that *Staphylococcus aureus* α-hemolysin channels in membranes tethered to gold have the same properties as those formed in free-standing bilayer lipid membranes. Neutron reflectometry provides high-resolution structural information on the interaction between the channel and the disordered membrane, validating predictions based on the channel’s x-ray crystal structure. The robust nature of the membrane enabled the precise localization of the protein within 1.1 Å. The channel’s extramembranous cap domain affects the lipid headgroup region and the alkyl chains in the outer membrane leaflet and significantly dehydrates the headgroups. The results suggest that this technique could be used to elucidate molecular details of the association of other proteins with membranes and may provide structural information on domain organization and stimuli-responsive reorganization for transmembrane proteins in membrane mimics.

INTRODUCTION

The primary goal of structural biology is to understand the structure-function relationship of proteins, which constitute the machinery of life. Despite the stunning achievements in determining protein structures using electron microscopy (1) and x-ray crystallography (2), the number of solved membrane protein structures significantly lags that of soluble proteins (3), in part because of the inherent difficulty of crystallizing proteins whose native environment is a disordered fluid membrane. Furthermore, these methods do not demonstrate whether the structural models are functionally accurate. NMR provides dynamic information about protein structure (4) but the technique is still limited to the study of relatively small proteins (i.e., with molecular mass <50 kDa) (5). Neutron reflectometry can provide complementary information to NMR and crystallography if a suitable interface for membrane protein reconstitution can be developed. The challenge lies in fabricating a biomimetic interface that retains the fluidity of a cell wall while providing long-term stability that is needed for structural characterization methods.

Tethered bilayer lipid membranes (tBLMs) combine the fluidity of a lipid bilayer with a stable platform for analysis with analytical techniques, including electrochemistry (6,7) and surface-sensitive scattering (8). Synthetic chemistry has recently been developed (8,9) that facilitates the formation of tBLMs in which the membranes are intrinsically disordered, thus permitting the reconstitution of proteins such as ion channels (10). Such membrane platforms are sufficiently resilient to be long-term stable (10,11) and can be characterized with neutron reflection at high resolution (8,12) by taking advantage of the possibility, offered by tBLM systems, to vary isotopic contrast and incorporate proteins in situ. This enables the acquisition of data sets before and after protein reconstitution, and under different isotopic labeling schemes, from one physical sample.

Although α-hemolysin (α-HL) spontaneously forms ion channels in free-standing bilayer lipid membranes (BLMs) (13), it is more difficult to reconstitute it into surface-bound membranes, presumably because of the rigidity of the membranes’ inner leaflets (14). With a novel synthetic approach to tBLM synthesis, successful reconstitution of the membrane channels was recently demonstrated (10). In this work, we prepared a mixed self-assembled monolayer (SAM) of a thiahexa(ethylene oxide)-substituted lipid analog, 20-tetradecyloxy-3,6,9,12,15,18,22-heptaoxaheptatriocantone-1-thiol (WC14) (8) and β-mercaptoethanol (β-ME), then completed the bilayer with phospholipid via rapid solvent exchange (see Methods). Electrochemical impedance spectroscopy (EIS) and neutron reflection (NR) show that the resultant tBLMs are impermeable to ions and contain water in the submembrane space (8), which should facilitate protein reconstitution (7,15).

MATERIALS AND METHODS

Materials

WC14 was synthesized, purified, and characterized in house, as reported previously in the Supplementary Material of McGillivray et al. (8).
1,2-diphtanoyl-sn-glycero-3-phosphatidylcholine (DPhyPC) and pyridyldithiopropionate-poly(ethylene glycol)-distearoyl-sn-glycerophosphatidylethanolamine (PDP-PEG2000-DsPE) were from Avanti Polar Lipids (Birmingham, AL). β-ME (Sigma-Aldrich, St. Louis, MO) was distilled before use. Poly(ethylene glycol) (PEG) and D2O (99.9% isotope purity) were obtained from Fluka (Ronkonkoma, NY) and Cambridge Isotopes Laboratory (Andover, MA), respectively. H2O was from a Millipore (Billerica, MA) UHQ reagent-grade water purification system.

Conductance measurements of α-HL in free-standing bilayer membranes

Freestanding BLMs (16) were formed from DPhyPC on ~100 μm diameter holes in a thin Teflon partition separating two identical Teflon chambers (17) that each held ~2 mL of electrolyte solution (0.1 M KCl, 5 mM 3-((N-morpholino)propanesulfonic acid (MOPS), 5 mM 2-(N-morpholino)ethanesulfonic acid (MES) at either pH 7.5 or 4.5). A small voltage (~2 mV) was applied across the membrane via two matched Ag/AgCl electrodes, and the ionic current was converted to voltage and amplified with an Axon Instruments 200B patch clamp amplifier (Molecular Devices, Sunnyvale, CA) in the voltage clamp mode. In the voltage clamp mode, the membrane was held at a fixed potential and an applied current is converted to a voltage drop across the membrane. The potential difference is then measured and the ionic current is calculated from Ohm's law.

Data analysis was performed in terms of homogeneous layer models, or “box” models (20), using the ga_refl software developed at the NIST Center for Neutron Research (21). The reflectivity due to a model neutron scattering profile was computed using an optical matrix based on Parratt’s recursion algorithm (22). Isotrophy structures with distinct isotropic contrast were fitted simultaneously by consistently refining the corresponding nSLD profiles that were parameterized in terms of the underlying molecular structures (23,24). The robustness of the BLMs permitted the characterization of the membrane at various solvent contrasts followed by protein reconstitution and further NR characterization of the reconstituted BLM with the same physical sample without removal from the reflectometer. With the substrate and membrane structures thus established, the structural changes due to protein incorporation could be studied in full detail. Although the data at the highest momentum transfer (Q, > 0.2 Å−1) do not appear to contribute information about distinctive structural features of the BLM with and without the protein (see Fig. 3 below and the Supporting Material), these data were included in the error-weighted fitting because they contribute information about the overall sample structure.

For a refined data analysis, the crystal structure of the α-HL channel (Protein Data Bank accession code: 7AHL) (25) was incorporated into the model by assuming that the barrel of the protein was aligned perpendicular to the membrane (z direction) and the crystal structure was undisturbed. The contribution of the protein to the nSLD profile along z was then divided into 0.5 Å thick slices in which the sums of the atomic scattering lengths, divided by the solvent-excluded volumes, were calculated. The solvent-excluded volume was determined by the method of Connolly (26), using a 1.4 Å radius probe to represent a water molecule. In the fitting to the data, the resultant protein nSLD contribution was allowed to shift freely along z relative to the bilayer membrane. The lateral protein density at/in the membrane was accounted for as a fitting parameter. Protein contributions in the aqueous volume phase outside the membrane slabs were integrated into six layers. Measurements were typically made in three solvent contrasts (D2O, H2O, and a D2O/H2O mixture with an nSLD of 4 × 10−6 g cm−3, “CM4”). All solvent contrasts were simultaneously fitted for the BLM with and without α-HL (21).

RESULTS AND DISCUSSION

Electrochemical investigation of biomimetic interface and of protein functionality

The addition of α-HL to the aqueous phase caused significant changes in the magnitude of the tBLM impedance, |Z|, and the phase angle (Fig. 1). These EIS features are associated with ion flux induced by membrane ionophores (6) . Quantitative information was obtained by modeling the
impedance spectra with the equivalent circuit shown in Fig. 1 (for details of the fitting, see the Supporting Material). The membrane is characterized by a constant phase element \( (\text{CPE})_{\text{tBLM}} \) \((27)\), with \( \alpha = 0.986 \pm 0.001 \) \((n = 22 \text{ samples})\), which suggests a nearly ideal capacitive behavior of the electrically insulating dielectric bilayer \((14)\). The \( \text{CPE}_{\text{tBLM}} \) coefficient, \( (0.62 \pm 0.01) \mu F \times cm^{-2} s^{(\alpha-1)} \) \((n = 22 \text{ samples})\), agrees with the specific capacitance values for solvent-free BLMs \((28)\). Upon incubation with \( \alpha \)-HL, \( \text{CPE}_{\text{tBLM}} \) increases in a protein concentration- and time-dependent manner, which suggests a net increase in the bilayer dielectric constant upon insertion of water-filled ion channels into the membrane \((29)\). In addition, the reconstitution of ion channels creates parallel conductance pathways that bypass the membrane’s high resistance. These pathways are represented by another constant phase element, \( \text{CPE}_{\text{pores}} \), in series with a resistor, \( R_{\text{pores}} \). The nearly ohmic open channel conductance determines \( R_{\text{pores}} \), whereas \( \text{CPE}_{\text{pores}} \) is determined by a complex combination of the conductance of the aqueous reservoir between the Au surface and the inner membrane and the differential capacitance of the Au/reservoir interface.

Several criteria were used to demonstrate that the \( \alpha \)-HL channels in the tBLMs were equivalent to those in free-standing BLMs. First, the toxin increased the tBLM conductance in proportion to both the bulk electrolyte conductivity and the ion (i.e., the conductance is greater in a KCl than in an NaCl solution), as it does in BLMs \((13)\) (data not shown). Second, the rates at which \( \alpha \)-HL increased the conductance of tBLMs correlated with those of free-standing BLMs at the same pH values (Fig. 2 A). Third, the conductance

**FIGURE 1** EIS spectra of a tBLM before and after \( \alpha \)-HL reconstitution. The Bode plots \((A)\) and phase angles \((B)\) were measured with tBLMs in 0.1 M NaCl, 5 mM phosphate buffer at pH 7.5 before \((\text{open circles})\) and 2 h after the addition of \( \alpha \)-HL to a final concentration of 50 nM \((\text{solid circles})\). The temperature was \((21 \pm 1)^{\circ} \text{C}\). The data were normalized with respect to the membrane surface area. \((\text{Inset})\) Equivalent model circuit for the tBLM.

**FIGURE 2** \((A)\) Kinetics of \( \alpha \)-HL channel formation in free-standing and tethered bilayer lipid membranes. The rate at which \( \alpha \)-HL increased the specific conductance of tBLMs was greater at pH 4.5 \((\text{up triangles})\) than at pH 7.5 \((\text{downward triangles})\). Similar results were obtained with free-standing BLMs \((\text{solid lines})\). The applied potential was 10 mV alternating current (tBLM) and 2 mV direct current (BLM). In the BLM experiments, \( \alpha \)-HL was added to the solution bathing one side of the membrane. In both experiments, the bulk concentration of \( \alpha \)-HL was 50 nM. For the EIS data, the model-derived conductance, \( Y_{\alpha \text{-HL}} = R_{\text{pores}}^{-1} \), was normalized with respect to the surface area. \((B)\) Effect of size-selected PEGs on \( \alpha \)-HL channel conductance. Polymers with molecular mass \( \% 2000 \text{ g/mol} \) decreased the conductance of tBLMs containing many \( \alpha \)-HL channels \((\text{open diamonds})\) and of single \( \alpha \)-HL channels in a free-standing BLM \((\text{solid diamonds})\) \((30)\). The concentration of PEG in each experiment was 15\% (w:w). The curves are drawn to guide the eye. The bottom dashed line indicates the ratio of the bulk conductivities, \( \sigma(\text{PEG})/\sigma(\text{PEG}) \).
increased superlinearly with the bulk $\alpha$-HL concentration in both membrane types (data not shown). In general, the specific conductance increases induced by $\alpha$-HL in tBLMs were within an order of magnitude (usually within a factor of 2 to 7) of those in free-standing BLMs.

The diameter of the $\alpha$-HL channel in free-standing BLMs was estimated previously from the effect of PEG on the channel conductance (30). The conductance decreases when the PEGs are small enough to partition into the pore. For $\alpha$-HL, this limit is $\sim 2250$ g/mol. Fig. 2 B shows that the effect of different molecular mass PEGs on the conductance of $\alpha$-HL-doped tBLMs (open symbols) is similar to that on single $\alpha$-HL channels in free-standing BLMs (solid symbols). The difference between the two partitioning plots might be due to restricted diffusion of PEG in the region adjacent to the Au electrode on the solid surface. Nevertheless, the results suggest that the values of the conductance, $Y_{\alpha$-HL} = R_{pores}^{-1}$, derived from the EIS model are directly related to the intrinsic conductance of $\alpha$-HL.

The correspondence between the results in BLMs and tBLMs demonstrates that $\alpha$-HL channels are functional in both systems. Thus, other techniques such as NR can be used to determine the structural characteristics of the $\alpha$-HL-membrane complex, taking advantage of the tBLM system’s high stability.

**Neutron reflectometry of the biomimetic interface**

The sample preparation led to a robust film that was stable for days and survived solvent exchanges for isomorphic contrast adjustments (12). The multiple solvent contrasts allow for unambiguous determination of model parameters. For details of the technique, see Majkrzak et al. (31) and references therein. Fig. 3 shows NR results for a tBLM with and without $\alpha$-HL in D$_2$O (for full data sets, see the Supporting Material). Changes in the scattering after exposure to $\alpha$-HL are shown as error-weighted residuals of reflectivity data against the best-fit reflectivity model of the protein-free tBLM. Deviations from the baseline are most pronounced at low momentum transfer (e.g., around $Q_z = 0.05$ Å$^{-1}$, where the coordinate $z$ is normal to the membrane plane and originates at the gold/thiol interface). This indicates significant structural changes of the bilayer upon $\alpha$-HL reconstitution on the length scale, $2\pi/Q_z \approx 120$ Å, the approximate height of the $\alpha$-HL channel along its symmetry axis (25). Although less pronounced than those around $Q_z = 0.05$ Å$^{-1}$, significant deviations persist to 0.2 Å$^{-1}$. However, more information can be extracted from these data sets by using more detailed fitting of the data, with additional details gleaned from the use of multiple solvent contrasts.

Simultaneously fitting the data in three solvent contrasts to a slab model with a single extramembranous layer accounts for the most prominent feature of the channel: its large cap domain exterior to the membrane (25). Structural details were derived from a composition-space model (23,24) of the membrane-inserted $\alpha$-HL array. This modeling procedure assumes the protein inserts into the membrane according to its known x-ray structure (25) and suggests that the nanopore spans the tBLM with its $\beta$-barrel stem end flush with the lipid headgroup region of the inner bilayer leaflet (Fig. 4, tabulated details can be found in the Supporting Material). Although this is not obvious from the nSLD profile (Fig. 4 A), the optimized parameter set determines the penetration depth to within $\pm 1.1$ Å, as shown in Fig. 5 A by a rigorous evaluation of parameter confidence limits with a Monte Carlo resampling technique (Supporting Material). The $\alpha$-HL stem is thus located $\sim 15$ Å away from the Au interface. Control experiments with $\alpha$-HL in a tBLM prepared with a longer tether (i.e., 60 Å) indicated that the proximity of the Au surface to the WC14-based tBLM does not alter the association of $\alpha$-HL with the membrane (Supporting Material).

A quantitative evaluation of the nSLD profile shows that the $\alpha$-HL channel reconstitutes into the tBLM at $\sim 33\%$ of...
the maximum hexagonal packing density, as set by the dimensions of the cap domain (25). The incorporation of such a large amount of protein without bilayer collapse is another indicator of the tBLM’s robustness.

In H$_2$O, a region of high nSLD around $z = 52$ Å (Fig. 4A) provides the location of the outer membrane headgroup layer. Incorporation of α-HL leads to a significant nSLD increase in this region, indicating a strong dehydration of the headgroups. The loss of hydration expected from the compression of the lipid bilayer by the α-HL stem would be on the order of 3%, based on the observed lateral density of the cap domain near the membrane and the protein geometry. NR shows that the α-HL channel removes most of the water from the lipid headgroups (Fig. 5B). This suggests that the protein cap interacts strongly with lipid headgroups and significantly perturbs their conformations, consistent with predictions based on the crystal structure (25). The overhang between the membrane-penetrating stem segment and the

![Supramolecular model of the tBLM reconstituted with α-HL.](image1)

FIGURE 4 Supramolecular model of the tBLM reconstituted with α-HL. Simultaneous fitting of a molecular model to five NR data sets (two for the tBLM and three for the protein-reconstituted tBLM) leads to five nSLD profiles that are all consistent with one supramolecular structure. (A) The three profiles for the tBLM reconstituted with α-HL. These profiles contain the calculated contribution of the α-HL x-ray crystal structure (21) at a lateral density and insertion depth derived from the model fit. (B) A scaled cartoon of the system.

![Parameter estimation and confidence limits for the supramolecular structure of α-HL reconstituted into a tBLM, determined with NR and quantified by Monte Carlo resampling (Supporting Material).](image2)

FIGURE 5 Parameter estimation and confidence limits for the supramolecular structure of α-HL reconstituted into a tBLM, determined with NR and quantified by Monte Carlo resampling (Supporting Material). Five individual data sets were linked in a composition-space modeling approach (23) that took advantage of the known α-HL crystal structure (21). Histograms of parameters describing the resampled data indicate that the protein can be localized along the surface normal with a precision of ± 1.1 Å within the fluid bilayer (A). (B) Both the lipid headgroups of the outer membrane leaflet (open symbols; dashed lines) and the content of the submembrane layer (solid symbols), including the membrane anchor and lipid headgroups of the inner leaflet, are significantly dehydrated upon reconstitution of α-HL. Parameter distributions for the submembrane layer hydration and protein insertion depth were approximately Gaussian (fitted lines). In contrast, the parameter distributions for the hydration of the outer lipid headgroup layer show extended tails and were evaluated by their mean values and standard deviations (for the full parameter set, see Table S3 in the Supporting Material).
cap domain is lined with hydrophilic residues that may form a suitable environment for zwitterionic phospholipid headgroups. Moreover, the edge of the cap contains hydrophobic residues that may interact with the acyl chains of the lipid bilayer. These results should aid Poisson-Nernst-Planck modeling of these systems because dehydration of the cleft between the cap domain and membrane will influence the dielectric properties of the channel-membrane system and could affect the channel conductance (32).

CONCLUSIONS

Although crystallography remains the technique of choice for determining protein structures with full atomic detail, a great deal of complementary information can be elucidated from disordered structures with other techniques, such as neutron reflectometry. Inspired by earlier work (7,18), we developed what to our knowledge is a new methodology for structural measurements on transmembrane proteins, verification of their functionality, and probing of their interactions with disordered membranes. The α-HL protein ion channel was chosen as a model system for proof-of-concept because its structure is known to high resolution (25). The technology we describe here will permit a broad range of biomedical investigations where the interaction of proteins with membranes is of immediate interest, such as studies into toxicology (33,34), Alzheimer’s disease (35), cell signaling involving lipids (36), or laminopathies (37).

Although the data interpretation described here utilized a known crystal structure, the techniques developed in this work can be readily extended to the use of other structural motifs as a basis. For instance, NR-derived structures of proteins with established functionality in a tBLM could be used in conjunction with computer models to discriminate between disparate solutions in the modeling. We anticipate that this technology will complement existing methods for both structure and functional measurements of membrane proteins.

SUPPORTING MATERIAL

More explanations and descriptions, figures, tables, and references are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(08)03231-1.

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REFERENCES


