Abstract: This paper proposes a method for sensing affinity interactions by triggering disruption of self-assembly of ion channel-forming peptides in planar lipid bilayers. It shows that the binding of a derivative of alamethicin carrying a covalently attached sulfonamide ligand to carbonic anhydrase II (CA II) resulted in the inhibition of ion channel conductance through the bilayer. We propose that the binding of the bulky CA II protein (MW ≈ 30 kD) to the ion channel-forming peptides (MW ≈ 2.5 kD) either reduced the tendency of these peptides to self-assemble into a pore or extracted them from the bilayer altogether. In both outcomes, the interactions between the protein and the ligand lead to a disruption of self-assembled pores. Addition of a competitive inhibitor, 4-carboxybenzenesulfonamide, to the solution released CA II from the alamethicin-sulfonamide conjugate and restored the current flow across the bilayer by allowing reassembly of the ion channels in the bilayer. Time-averaged recordings of the current over discrete time intervals made it possible to quantify this monovalent ligand binding interaction. This method gave a dissociation constant of ~2 μM for the binding of CA II to alamethicin-sulfonamide in the bilayer recording chamber: this value is consistent with a value obtained independently with CA II and a related sulfonamide derivative by isothermal titration calorimetry.

Introduction

This paper describes a method to quantify protein–ligand interactions by measuring the flux of ions through pores of a synthetically modified, ion channel-forming peptide in planar lipid bilayers. The method is based on the reduction of ionic conductivity through self-assembled pores of a derivative of alamethicin, a naturally occurring ion channel-forming peptide,1–3 which carries a covalently attached benzenesulfonamide group on its C-terminus (alamethicin sulfonamide, 8, see Scheme 1). In this assay, reduction of the ion current is triggered by binding of carbonic anhydrase II (CA II) to the sulfonamide moiety on the channel-forming peptide. By time-averaging the recorded current, we were able to compare the amount of transported charge before and after addition of CA II. Increasing concentrations of CA II reduced ion transport. This reduction made it possible to estimate binding constants for the interaction between CA II and the sulfonamide ligand (by plotting transported charge through the pores as a function of the concentration of CA II). Addition of a competitive inhibitor, 4-carboxybenzenesulfonamide (9), to the solution released CA II from the complex with 8 and restored the current flow through self-assembled ion channels in the bilayer.

Ion channels are increasingly being investigated for sensing applications,4–11 because binding of just one ligand molecule can induce a conformational change of the ion-channel protein and result in the flux of thousands of ions. In this sense, ion channels are amplifiers and can amplify signals by factors of 103–105.12 This high-gain amplification makes ion channels well suited for signal transduction, which is, of course, one of their critical functions in the membranes of excitable cells.13

On the basis of pioneering work on protein engineering by the groups of Mutter,14–17 Montal,18,19 DeGrado,20,21 Vogel,22–25

Using Ion Channel-Forming Peptides to Quantify Protein–Ligand Interactions
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Synthesis of an Alamethicin Derivative Carrying a Benzenesulfonamide Group

\[
\begin{align*}
&\text{Ac-Alb-Pro-Alb-Alb-Alb-Gln-Alb-Val-Alb-} \\
&\quad \text{Gly-Leu-Alb-Pro-Alb-Alb-Glu-Gln-} \\
&\quad \text{alamethicin, 6}
\end{align*}
\]

4-carboxybenzenesulfonamide, 9

Reagents and Conditions: (a) (Boc-aminooxy)acetic acid, EDE, DIEA, DMF (b) Pd/C, H\(_2\), EtOH (c) I, EDC, DIEA, DMF (d) TFA (e) Dess-Martin oxidation, CH\(_2\)Cl\(_2\) (f) 5, CH\(_2\)OH.

and Bayley,\(^{26-30}\) ion-channel proteins and ion-channel-forming peptides have received increasing attention for sensing applications since the 1990s.\(^{31}\) In 1993, Rokitskaya et al. explored photodynamic inactivation of gramicidin pores in the presence of phthalocyanines.\(^{32}\) One year later, Bayley’s group demonstrated the use of \(\alpha\)-hemolysin for detecting metal ions\(^{36}\) and for sensing protease activity,\(^{33}\) whereas Bezrukov et al. used alamethicin pores to count polymer particles.\(^{34}\) Woolley et al. used a derivative of alamethicin to form an “ion-activated” ion channel.\(^{35}\) Cornell et al. employed gramicidin derivatives for detection of immune reactions;\(^{36}\) and Vogel’s group developed a synthetic ion channel\(^{23}\) and detected binding of colicin N to the porin OmpF.\(^{24}\) Since these initial studies, melittin,\(^{4,23,25,37,38}\) gramicidin A,\(^{5,41}\) melittin,\(^{338}\) \(\alpha\)-hemolysin,\(^{27,39,40}\) gramicidin A,\(^{54-47}\) and alamethicin\(^{36,34,48-56}\) have become the most extensively studied systems for ion...
channel-based sensor development. The reasons for the interest in these four systems are that: (i) they incorporate spontaneously into bilayers from solution (in contrast to most ion-channel proteins, which must be reconstituted into the bilayer by techniques such as proteoliposome fusion);57 and (ii) they are available commercially.

In a series of elegant studies, Bayley and co-workers showed that wild-type or genetically modified α-hemolysin pores can be used to detect individual molecules, binding interactions, reversible chemical reactions, or point mutations in DNA strands.39,58–61 Despite these scientifically compelling results, the use of α-hemolysin remains limited to specialized research laboratories. Two factors impede its widespread use: (i) experiments involving planar lipid bilayers require substantial technical expertise, and the bilayers are typically only stable for hours; (ii) rationally designed modification of α-hemolysin, using methods pioneered by Bayley, requires the tools and expertise of molecular biology. For a broader application of these systems, it would be useful if ion channel-forming peptides could be synthesized chemically at low cost, with high yield, high purity, and good stability; channel synthesis of (relatively) low molecular weight species would make it possible to produce variants at will, and to imagine certain large-volume applications that would be difficult for proteins.

Because they are relatively small molecules (<2 kD) and commercially available, melittin, gramicidin A, and alamethicin62 might become interesting building blocks for ion-channel sensors if they could be modified synthetically. Woolley et al. demonstrated, for instance, that the ion permeability through chemically modified gramicidin pores in a lipid bilayer could be used as a pH sensor.42 In a recent proof of principle study, the groups of Mayer and Yang showed that gramicidin A can be engineered to detect specific chemically reactive agents in solution.53,54 Futaki’s group employed peptide synthesis to prepare an analogue of alamethicin with a carboxylic acid group on the C-terminal end that could be modified selectively.53–55

these authors attached a biotin group to alamethicin and demonstrated binding of streptavidin and anti-biotin antibodies.51

Here we explored alamethicin as the platform for a sensor with three goals in mind: (i) to prepare derivatives of alamethicin by chemical derivatization of commercially available alamethicin that would make this type of sensor broadly available, (ii) to explore if sensors based on ion channel-forming peptides could be used to quantify protein–ligand interactions in solution, and (iii) to investigate a “typical”, biologically relevant, monovalent interaction between a protein and a small ligand. To accomplish these goals, we chose the well-characterized interaction between CA II (E.C. 4.2.1.1)64,65 and a benzenesulfonamide moiety. This interaction has a typical dissociation constant for biochemical interactions (~1 μM).66,67

Alamethicin is an antimicrobial peptide composed of 19 amino acids and one amino alcohol with a molecular weight of 1.96 kD. It is secreted by the fungus Trichoderma viride.68 Alamethicin adopts an amphiphatic, α-helical structure in biological membranes and forms ion channels by self-assembly to oligomers. The most accepted model of pore formation by alamethicin, the so-called barrel-stave model, suggests an arrangement of transmembrane helices in a circle with a central, water-filled pore.1,69 In this model, the hydrophilic face on the α-helix of alamethicin is oriented toward the lumen of the pore, whereas the hydrophobic face on the helix is in contact with the surrounding lipid molecules.69

Alamethicin monomers in aqueous solution bind to (or dissolve in) lipid membranes with partition coefficients of ~10−3 M. This value results in an equilibrium distribution of alamethicin in solution and alamethicin bound to the membrane (Figure 1).70 It also implies that it is relatively easy energetically to extract alamethicin from a lipid bilayer in which it is dissolved. Upon application of a transmembrane voltage, membrane-associated alamethicin molecules can adopt a transmembrane configuration in which the axis of the α-helix is oriented perpendicular to the plane of the bilayer (alamethicin has a permanent dipole moment along this axis, which corresponds to a net +0.5 charge at the N-terminus of the helix and a net −0.5 charge at its C-terminus).70 The probability for alamethicin monomers to adopt this perpendicular orientation increases strongly (nonlinearly) if the applied transmembrane voltage exceeds a certain threshold voltage. Once alamethicin adopts the transmembrane configuration, self-assembly of alamethicin monomers leads to pores that can comprise up to 11 monomers.3 This number of monomers in a pore fluctuates dynamically; these fluctuations lead to the characteristic stepwise changes between discrete conductance levels of single alamethicin pores.49,50,71 To perform single-channel recordings of alamethicin pores, a constant voltage above the threshold voltage has to be applied (in the work presented here, we used a

transmembrane voltage of 0.14 V for all experiments\(^{49}\). To record ion flux through a single alamethicin pore, the concentrations of the peptide, \([\text{alamethicin}]\), had to be low (\(\sim 5 \text{nM} \)) under the experimental conditions used in this work to avoid the occurrence of multiple pores at the same time.\(^{49,76}\) Increasing the concentration of the peptide, while keeping the transmembrane potential constant above threshold, led to an increase of alamethicin monomers in the transmembrane configuration and thus to a strong increase in the probability of channel formation due to the cooperative nature of pore formation.\(^{62}\) The resulting “macroscopic currents” across the membrane were due to the opening of multiple (often many) pores at the same time. If we kept all other experimental parameters constant (applied voltage, ionic strength, conductance of the electrolyte, lipid composition, bilayer thickness, surface area of the planar bilayer, etc.), then the “macroscopic conductance”, \(G\) (in \(\Omega^{-1}\) or Siemens, S), through a membrane with many alamethicin pores followed the relationship:

\[
G \propto [\text{alamethicin}]^n
\]

where \(n\), the power dependence of \(G\) on the peptide concentration, has previously been used as an estimate of the average number of alamethicin monomers in the channels. This simple approach (eq 1) provides, however, at best, only a rough estimate of the average number of monomers in the channels.\(^{73}\)

**Experimental Design.** We chose alamethicin for this study because we wanted to explore the possibility of interfering with a self-assembly process in such a way that it could be monitored in real time and, hence, exploited for sensing. Six characteristics of alamethicin make this ion channel-forming peptide an interesting choice for detecting the modulation of self-assembly processes. (i) Alamethicin pores are formed by the dynamic self-assembly of up to eleven monomers (in contrast to gramicidin pores that form by self-assembly of two peptides or melittin pores that form by self-assembly of four peptides); the variability of the composition of the pores results in a cooperative mechanism of pore formation and hence in a power-law of the dependence of the current through alamethicin pores on the concentration of alamethicin (eq I). This strong (nonlinear) response of the macroscopic current through alamethicin pores as a function of the alamethicin concentration is attractive for sensing, if the reaction that is used for detection can modulate this concentration and thus yield a strong response from the sensor system. (ii) The self-assembly of alamethicin can be monitored with high precision and with a time-resolution below milliseconds, by using established planar bilayer recording procedures.\(^{49,71}\) (iii) Alamethicin is commercially available and can be modified chemically, while preserving its ion channel-forming properties.\(^{51,52,74}\) The discrete conductance levels observed from single-channel recordings of alamethicin make it possible to examine the effect of chemical modification on the characteristics of the peptide that influence its ability to form ion channels.\(^{75}\) (iv) Alamethicin is a relatively small molecule, and thus, binding of a protein with a 10-fold larger molecular weight may be able to interfere with its self-assembly. (v) Alamethicin incorporates spontaneously into bilayers; spontaneous self-assembly makes its use more practical than possible uses of most ion-channel proteins. (vi) Alamethicin (or \(8\) in the work presented here), once added to the aqueous solution of a planar bilayer experiment, establishes a chain of equilibria (Figure 1) between free \(8\) in solution, \(8\) bound to the lipid bilayer, and \(8\) in an open, conducting pore. Assuming that molecules of \(8\) that are bound to CA II cannot participate in the formation of an open, conductive ion channel (either by inhibition of self-assembly of the complex of CA II–8 to a pore or by removal of CA II–8 from the bilayer), we hypothesized that reducing any one of the concentrations of \(8_{\text{solution}}, 8_{\text{bilayer}},\) or \(8_{\text{pore}}\) by binding to CA II will shift these equilibria. This shift will ultimately lower the concentration of alamethicin in open pores, \([8_{\text{pore}}]\). Due to the power law in eq 1, this reduction of \([8_{\text{pore}}]\) would lead to a strong reduction in the recorded transmembrane current.

For sensing applications, the stability of planar lipid bilayers is often a limiting factor. In order to generate lipid bilayers that were stable over several hours, we prepared planar membranes over micropores in Teflon films by the “folding technique” (a technique that prepares bilayers by apposition of lipid monolayers which had previously been spread at an air–water interface) for all experiments.\(^{49,76,77}\) These folded bilayers have the advantage that they form readily over small apertures (here, \(<50 \mu\text{m}\)), which increases the mechanical stability and reduces the electrical noise. Moreover, folded bilayers result in so-called “solvent-free” membranes, i.e., in lipid bilayers that contain only a small amount of organic solvent.\(^{76}\) We expected that the reduced amount of solvent would minimize fluctuations in bilayer area and thus facilitate reliable quantification of the current flux across the bilayer.\(^{77}\)
Results and Discussion

Synthesis. To make sensors based on ion channel-forming peptides practical, a straightforward synthesis of the sensing element that gives the required molecule (and variants of it) in good yield and high purity is desirable. Our work constitutes a step in that direction but is not yet a complete success. Several groups have reported methods of derivatization of alamethicin through C-terminal modification of native or synthetic alamethicin. The most common synthetic strategy that produces alamethicin derivatives with good to moderate ion channel activity is the formation of an ester or carbamate on the C-terminal phenylalaninol alcohol moiety of alamethicin. Our own attempts to make derivatives of alamethicin through esterification of the C-terminal alcohol resulted in impractically low yields and prompted us to explore new routes to derivatization of the C-terminus of alamethicin. Using HPLC-purified, commercially available alamethicin, we demonstrated that the C-terminal alcohol of alamethicin can be oxidized under mild conditions to the corresponding aldehyde in 50% isolated yield (Scheme 1). Reaction of this aldehyde with an O-alkylated hydroxyl amine carrying a benzenesulfonamide moiety afforded oxime (alamethicin sulfonamide, ) in 48% isolated yield. Although the strategy presented in Scheme 1 is not ideal due to the requirement for HPLC purification, it is, we believe, a first step toward a practical synthesis of derivatives of alamethicin useful in the applications we envision.

Ion-Channel Measurements. To characterize the ion channel activity of , we performed single-channel recordings in a planar bilayer setup (see Experimental Section for details), and compared the results to native alamethicin ( in Scheme 1). Figure 2 shows a representative trace of current versus time for both molecules. Interestingly, was capable of forming well-defined conductance states, although its single channel conductance was slightly reduced compared to native alamethicin (conductance states or conductance levels refer to assemblies of alamethicin that formed pores with discrete conductance of ions through the bilayer). This reduction was most pronounced at small conductance levels: conductance level 1 of reached 52% of conductance level 1 of native alamethicin, conductance level 2 reached 61%, conductance level 3 reached 73%, and conductance level 4 reached 78% of the conductance of the corresponding level of native alamethicin. In addition, the derivatized molecule resulted in noisier traces than native alamethicin.
This additional noise may be caused by structural fluctuations of the covalently attached sulfonamide group\textsuperscript{85} (i.e., the attached group may fold into and out of the pore) or by small-scale dynamic rearrangements of the self-assembled pore due to the presence of the sulfonamide moiety. The observation that the difference in conductance between \textsuperscript{8} and native alamethicin was most pronounced in small pores supports the hypothesis that the sulfonamide moiety partly occluded the channel by folding toward or into it; Figure 3 illustrates schematically that such an occlusion would have a larger effect on small pores than on large pores. Figure 3B and C shows that the size of the sulfonamide group with its linker (5) relative to the size of alamethicin (6) may affect the ion channel conduction through pores of \textsuperscript{8} if 5 were to bend toward the lumen of the pore. The energy-minimized structures in Figure 3B and C represent only one plausible conformation of \textsuperscript{8}; the flexible backbone of 5 is expected to adopt a number of different conformations in solution.

Figure 4 illustrates the concept of an affinity sensor based on ion channel-forming peptides. The ionic conductance through a pore is modulated by specific binding of a macromolecule to the peptides that form the pore. Futaki’s group proposed that in their experimental system of biotinylated alamethicin, the reduction in current was due to blocking of alamethicin pores in a planar lipid bilayer. The current (flux of ions) through single or multiple pores is recorded with a patch clamp amplifier. (B) Addition of a macromolecule (here CA II, represented in blue) that binds to the ligand, may have two consequences: (1) it may disrupt the pore, either by steric hindrance, or by removing the peptide from the bilayer (see asterisk), or (2) it may bind close to the mouth of the pore and block it. In either case, only a small (or no) current would be recorded as a result of the interaction. (C) Addition of competitive ligand (here 9, small gray arrows) to the solution leads to binding of free ligand to the proteins, and releases ion channel-forming peptides. This action makes it possible for the peptides to self-assemble again to a conducting pore, or it unblocks an existing pore that was blocked by the protein.

benzenesulfonamide group on this peptide (Futaki’s group had mentioned this mechanism as a possibility).\textsuperscript{51} This mechanism of action, if correct, could lead to complete cessation of ion channel activity, given that the concentration of CA II is sufficiently high that the majority of \textsuperscript{8} is bound to CA II (and that no impurities of 6 are present). Another mechanism for impeding the assembly of the pores might be extraction of monomers of \textsuperscript{8} entirely from the bilayer by formation of a soluble CA II-8 complex; this complex may be less prone to membrane interactions than free \textsuperscript{8}. It is also possible that several mechanisms, including blockage of pores,\textsuperscript{51} act in parallel or sequentially, depending on the concentration of CA II and \textsuperscript{8}.

Quantitative Analysis of Transported Charge through Bilayers by Time-Averaging. In the work presented here, we performed planar lipid bilayer experiments to record macroscopic currents through pores from an alamethicin derivative, \textsuperscript{8}. To determine the dependence of the recorded current, \( I \), on the concentration of \textsuperscript{8}, we analyzed the ion channel current over intervals of time \( t \) of 5 min. The area under the resulting current–time trace corresponded to the transported charge over the 5 min interval, \( Q_t \):

\[
Q_t = \int_{0}^{5} I(t) \, dt
\]

We then used this time-averaging approach to quantify the binding of CA II to \textsuperscript{8} based on the hypothesis that the binding of the comparably bulky CA II protein (molecular weight \( \sim 30 \) kD) to the comparably small peptide \textsuperscript{8} (molecular weight \( \sim 2.5 \) kD) would impede the bound molecules of \textsuperscript{8} from participating in functional self-assembly to a conducting pore. By comparing
the value of the transported charge during an interval of 5 min, $Q_S$, before and after incubating a constant concentration of 8 with increasing concentrations of CA II, we explored the possibility of determining the binding constant between CA II and 8.

Figure 5A and B illustrates this time-averaging approach to the binding of CA II to 8. According to eq 2, we integrated the area under the curve of current versus time (illustrated by the yellow area in Figure 5A) to obtain the total charge $Q_S$ (in Coulomb, C) that was transported through the bilayer over an interval of 5 min. Figure 5 A shows that a concentration of 40 ng mL$^{-1}$ (or 16.1 nM) of 8 in the bilayer chamber resulted in macroscopic currents that could exceed 3 nA. Because the amplifier with the gain setting that we used cannot measure currents above 20 nA due to current overload, we chose, somewhat arbitrarily, the concentration of 16.1 nM of 8 as the maximum concentration that would ensure, under all conditions, that even rare current spikes with large amplitude could be recorded accurately by the amplifier. We termed the charge that was transported during 5 min through bilayers in the presence of 16.1 nM 8, the maximum transported charge, $Q_{5\text{max}}$. Because our goal was to establish a repeatable method for quantifying the binding of CA II to 8 based on the measured current through pores of 8, we had to account for variations in the surface area of the planar lipid bilayers from experiment to experiment (the solvent torus that surrounds planar bilayers can vary in thickness and area even if the same aperture is used in repeated experiments$^{76,86}$). To account for these variations in bilayer area, we determined first the value of $Q_{S\text{max}}$ by adding 16.1 nM 8 to both bilayer chambers before starting each binding experiment. Then, after adding for instance CA II, we normalized the recorded transported charge $Q_S$ over the particular value of $Q_{S\text{max}}$ that was determined just before addition of CA II and termed the resulting normalized charge, “relative transported charge, $Q_{5\text{r}}$”.

Relative transported charge over 5 min:

$$Q_{5\text{r}} (\text{in } \%) = \frac{Q_S}{Q_{5\text{max}}} \times 100$$

(3)

Figure 6 shows that $Q_{5\text{r}}$ depended strongly on the concentration of 8 in the recording chambers. The increase in $Q_{5\text{r}}$ (in %) was fitted well ($R^2 = 0.999, N = 7$) with an equation of the type:

$$Q_{5\text{r}} = 3.4 \times 10^{35} \times [8]^{1.6}$$

(4)

We chose a power law based on eq 1, because current and thus transported charge are proportional to the conductance $G$. In eq 4, the concentration of 8 is expressed in units of M (i.e., mol L$^{-1}$) and the constant 3.4$ \times 10^{35}$ is expressed in units of M$^{-1.5}$.

**Determination of Equilibrium Binding Constants.** Figure 6B shows the effect of the addition of CA II to the compartments of the recording chamber containing 16.1 nM 8. The channel activity showed a strong reduction in ionic conductivity and hence in $Q_{5\text{r}}$ upon increasing the concentration of CA II from 0 to 2 $\mu$M. We suggest that this reduction in trans-bilayer current was due either to disruption of the self-assembled pores, or to blocking of the pores by binding of CA II close to the mouth of the pore (Figure 4). The line in Figure 6B represents a best fit to a model that is based on binding of CA II to 8. In this model, we made the following nine assumptions: (i) Only “free” molecules of 8 (i.e., 8 that was not bound to CA II) could contribute to the formation of conductive alamethicin pores. This assumption does not imply that, at any given time, all free molecules of 8 actually participated in forming a pore, only the molecules of 8 that were located in the bilayer and self-assembled to pores participated in conducting channels, but it does imply that only free molecules of 8 could participate in the chain of equilibria that could lead to open pores (illustrated in Figure 1). Binding of 8 to CA II would thus effectively reduce the concentration of free 8 and consequently shift the equilibria in Figure 1 in such a way that less free 8 would be available in the bilayer to participate in the formation of conductive pores than before the addition of CA II. (ii) Binding of one molecule of CA II “removed” only one molecule of 8, it did not remove aggregated clusters involving multiple molecules of 8. In contrast to this assumption, if a single molecule of CA II would block a conducting pore, then binding of each molecule of CA II would “remove” at least three molecules of 8 since the smallest conducting alamethicin pores are believed to consist of at least three (probably four) alamethicin monomers.$^{87}$ The model for quantification employed here is hence different from


either free or bound to CA II; only one affinity state existed and no partial binding occurred. (vii) Binding did not alter 8 or CA II irreversibly. (viii) Binding of 8 to CA II was the same regardless of the location where binding occurred (i.e., in solution, in the bilayer, or in a pore).

The fit to a power law in Figure 6A yielded an empirical relationship between \( Q_5 \) and the total concentration of 8 in the recording chambers. Because in Figure 6A the total concentration of 8 was equal to the free concentration of 8 (because no CA II was present and hence 8 could not be bound to CA II), we were able to use the measured values of \( Q_5 \) in Figure 6B to calculate the concentration of free 8 at each concentration of CA II in the reaction mixture. With this information, it is possible to calculate the concentration of 8 that is bound to CA II and hence to determine a binding constant by fitting the data in Figure 6B. To derive the function to fit the data in Figure 6B, we used the law of mass action for the binding reaction between CA II and 8:

\[
\text{CA II} + 8 \rightarrow \text{CA II-8}
\]

where \( K_{a8} \) represents the affinity constant of the binding reaction shown in eq 5. [CA II-8]_eq is the molar concentration of the complex of CA II with 8 at equilibrium, [8]_eq is the molar concentration of free 8 at equilibrium, and [CA II]_eq is the molar concentration of free CA II at equilibrium. Because we ultimately wanted to fit the data in Figure 6B, we were seeking a functional relationship between \( Q_5 \) and the concentration of CA II that was initially added to the recording chambers, [CA II]_0 (i.e., the concentration of CA II before binding to 8). As mentioned previously, eq 4 gave a relationship between \( Q_5 \) and [8]_eq. Because we assumed that only free 8 (i.e., not 8 bound to CA II) could participate in pore formation, the concentration of 8, [8]_eq, in eq 4 referred, in this model, to the concentration of free, unbound 8 at equilibrium, [8]_eq. Hence, the relationship between \( Q_5 \) and [8]_eq can be expressed as:

\[
Q_5 = 3.4 \times 10^{35} [8]_eq^{4.3}
\]

Rearranging eq 6 yields:

\[
[8]_eq = \frac{[\text{CA II-8}]_eq}{K_{a8} \cdot [\text{CA II}]_eq}
\]

Conservation of mass requires that eqs 9 and 10 hold ([8]_0 = 16.1 \times 10^{-9} M):

\[
[\text{CA II-8}]_eq = [8]_0 - [8]_eq
\]

\[
[\text{CA II}]_eq = [\text{CA II}]_0 - [\text{CA II-8}]_eq
\]

Because the total concentration of 8, [8]_0, in the recording chambers was low (nanomolar range) compared to the concentrations of [CA II]_0 that we added to the recording chambers (micromolar range), we assumed that the concentration of free CA II in equilibrium would be approximately the same as the initial concentration of CA II:

\[
[\text{CA II}]_eq \approx [\text{CA II}]_0
\]
Incorporating eq 9 and 11 into eq 8 yields:

$$[8]_{\text{eq}} = \frac{[8]_0 - [8]_{\text{eq}}}{K_{a8}[\text{CA II}]_0}$$  \hspace{1cm} (12)

and solving eq 12 for $[8]_{\text{eq}}$ gives:

$$[8]_{\text{eq}} = \frac{[8]_0}{K_{a8}[\text{CA II}]_0 + 1}$$  \hspace{1cm} (13)

Finally, combining eq 13 with eq 7, gives the desired relationship between $Q_{S\text{r}}$ and the concentration of CA II that we added to the recording chambers, $[\text{CA II}]_0$:

$$Q_{S\text{r}} = 3.4 \times 10^{-35} \left( \frac{[8]_0}{K_{a8}[\text{CA II}]_0 + 1} \right)^{1.3}$$  \hspace{1cm} (14)

We used eq 14 to fit the data in Figure 6B and obtained $K_{a8}$ as the only fitting parameter. We performed this analysis twice; the first fit included all points in Figure 6B (dotted line). This curve fit returned a value of $K_{a8} = (3.6 \pm 1.5) \times 10^5 \text{ M}^{-1}$ ($R^2 = 0.742$, $N = 8$) corresponding to a $K_{a8} = 2.8 \mu\text{M}$. Because the datum that is marked with an asterisk in Figure 6B appears to be an outlier (Figure 6B inset), we also performed the best fit analysis by excluding this point from the analysis (solid line). This approach returned a best fit with a value of $K_{a8} = (6.7 \pm 1.5) \times 10^2 \text{ M}^{-1}$ ($R^2 = 0.963$, $N = 7$), corresponding to a dissociation constant, $K_{a8} = K_{a8}^{-1} = 1.5 \mu\text{M}$.\(^{90}\)

To compare these values with a well-established method for determining binding constants, we transformed the data in Figure 6B to construct a binding isotherm (see Figure S1, Supporting Information). From the best curve fit analysis of this binding isotherm, we obtained a value of $K_{a8} = 0.9 \pm 0.2 \mu\text{M}$ when we excluded the apparent outlier (marked with an asterisk in Figure 6B and Figure S1) and a value of $K_{a8} = 1.3 \pm 0.3 \mu\text{M}$ when we included the outlier in the best fit analysis. In summary, depending on the transformation of the original data and the corresponding best curve fit analysis, we obtained a mean value of $K_{a8}$ ranging from 0.9 to 2.8 $\mu\text{M}$ for the interaction between $\mathbf{8}$ and CA II in the recording chambers.

For comparison with an independent method, we measured the affinity constant of CA II with benzylsulfonylamide (5) by isothermal titration calorimetry (ITC).\(^{91}\) We used precursor 5 for the ITC measurement because it was more soluble in aqueous solutions than 8; this increased solubility was required to achieve detectable reagent concentrations for ITC measurements. The resulting dissociation constant $K_{a8}$ by ITC was $2.6 \pm 0.6 \mu\text{M}$.


\(^{90}\) We tested the validity of the simplification that we made in eq 11 by performing the same analysis and fit of the data in Figure 6B without any simplification. In this case, we used eq 10 to obtain $[\text{CA II}]_{\text{eq}}$, which then lead to a quadratic equation to solve for $[8]_{\text{eq}}$ of the form: $[8]_{\text{eq}} = (1 - K_{a8}[\text{CA II}]_0 + K_{a8}[\text{CA II}]_0^2) \pm (1 - K_{a8}[\text{CA II}]_0 + K_{a8}[\text{CA II}]_0^2 + 2K_{a8}[8]_{\text{eq}})^{1/2}$. Using this precise analysis, we obtained $K_{a8} = (6.71 \pm 1.53) \times 10^2 \text{ M}^{-1}$ and, hence, the same mean value for $K_{a8}$ as the one we obtained by using the simplification in eq 11.


Figure 7. Relative transported charge $Q_{S\text{r}}$ as a function of increasing concentration of competitive inhibitor $\mathbf{9}$ in both compartments of the recording chamber. Both chambers initially contained $[\mathbf{8}]_0 = 16.1 \text{ nM}$ and $[\text{CA II}]_0 = 9.3 \mu\text{M}$. The solid curve represents a sigmoidal best fit to eq 15. This function approaches $Q_{S\text{r}} = 100\%$ for large concentrations of $[\mathbf{9}]_0$, as expected in the presence of an excess of competitor $\mathbf{9}$, which would release all $\mathbf{8}$ from the complex with CA II ($K_{a8} = 0.73 \pm 0.02 \mu\text{M}^{-1}$). The data shown as round symbols represent the fraction of $\mathbf{8}$ bound ($\%\text{bound}$) to CA II as a function of $[\mathbf{9}]_0$. The values for $\%\text{bound}$ were calculated from $Q_{S\text{r}}$ for every value of $[\mathbf{9}]_0$ using eqs 7 and 16. The $IC_{50}$ value for the displacement of $\mathbf{8}$ from CA II by adding increasing concentrations of competitive ligand $\mathbf{9}$ was determined with a best fit analysis of the round points with eq 17 (dotted curve). Note the logarithmic scale for $[\mathbf{9}]_0$. As recommended by Motulsky and Christopoulos, we chose, somewhat arbitrarily, log[(9)]0 = $-9$ for [9]0 = 0 $\mu\text{M}$, because log(0) is not defined, the corresponding data are marked by open symbols. The data in this figure were recorded only once: hence the absence of error bars.

Comparing binding well with the range of 0.9–2.8 $\mu\text{M}$ obtained with the ion channel system presented here.

**Competitive Binding Assay: Displacement of 8 from CA II.** To test if it was possible to displace the bound molecules of $\mathbf{8}$ from CA II by competitive binding, we added increasing concentrations of a competitive inhibitor, $\mathbf{9}$ (Scheme 1), to both compartments of a recording chamber which contained a total concentration of 16.1 nM of $\mathbf{8}$ and 9.3 $\mu\text{M}$ of CA II (i.e., $[\mathbf{8}]_0 = 16.1 \text{ nM}$; $[\text{CA II}]_0 = 9.3 \mu\text{M}$). We expected 9 to be able to displace $\mathbf{8}$ from CA II because the dissociation constant for binding of $\mathbf{9}$ to CA II, $K_{a9}$, is 0.73 ± 0.02 $\mu\text{M}$.\(^{67}\) Figure 7 (data shown as squares) shows that increasing concentrations of $\mathbf{9}$ resulted in an increase in the relative transported charge $Q_{S\text{r}}$ through the bilayer. This increase corresponded to an increased concentration of free $\mathbf{8}$ in solution as expected if $\mathbf{9}$ displaced $\mathbf{8}$ from the CA II-8 complex. The square-shaped points in Figure 7 could be fitted well ($R^2 = 0.999$, $N = 4$) with a dose–response curve (solid curve) of the following form:\(^{93}\)

$$y = y_{\text{min}} + \frac{y_{\text{max}} - y_{\text{min}}}{1 + 10^{\log EC_{50} - y \text{HillSlope}}} \hspace{1cm} (15)$$

where $y$ corresponds to $Q_{S\text{r}}$, $y_{\text{min}}$ represents the lower asymptotic value of the sigmoidal function, $y_{\text{max}}$ represents the upper asymptotic value, log $EC_{50}$ represents the logarithm of the concentration that generates 50% response ($EC_{50}$); $x$ corresponds to log([9])0, and HillSlope determines the steepness of the curve. Before fitting the data (squares) in Figure 7 to eq 15, we set $y_{\text{min}}$ to 0.1% because we measured this value of $Q_{S\text{r}}$ experi-
mentally before adding 9 (i.e., [9]₀ = 0) and we set y_{max} to 100% because we assumed that at high concentrations of competitive ligand (here [9]₀ > 1 mM) virtually all of 8 would be released from the CA II-8 complex (i.e., [8]_{eq} ≈ [8]₀ = 16.1 nM) and that consequently the relative transported charge Q_{35} would reach 100%. The best fit analysis of the square points returned a value for log IC₅₀ of −4.62 ± 0.01, corresponding to EC₅₀ = 24.2 µM and a value for the HillSlope of 4.51 ± 0.04.

To determine the dissociation constant K₉₉ for the interaction between CA II and 9 based on the competitive binding data in Figure 7, we used an approach developed by Linden in 1982. This approach makes it possible to calculate the dissociation constant from competitive binding curves without the requirement that the concentration of any of the involved species has to remain constant (i.e., it allows for depletion of all species). We used this approach for accurate determination of K₉₉ because we expected that the addition of micromolar concentrations of 9 to the recording chambers which contained nanomolar concentrations of 8 and micromolar concentrations of CA II would lead to significant changes in the equilibrium concentrations of free molecules of 8, 9, and CA II. Simplifications such as the one made in eq 11 would not be appropriate for determining K₉₉ in this case.

Application of Linden’s approach proceeded in three steps. First we used the competitive binding data in Figure 7 to determine the “inhibitory” concentration of 9 that displaced 50% of 8 from CA II, i.e., the IC₅₀, competition for competitive binding of 9 to CA II in the presence of 8 (Figure 7). We determined the IC₅₀ value by plotting $\Theta_9$-bound which is defined as

\[ \Theta_9^{\text{bound}} = \frac{[\text{CA II} - 8]_{\text{eq}}}{[8]_0} = \frac{[9]_0 - [8]_{\text{eq}}}{[9]_0} \]  

(16)

as a function of [9]₀ (round symbols in Figure 7) followed by fitting the data with an equation for competitive binding:

\[ y = y_{\min} + \frac{y_{\max} - y_{\min}}{1 + 10^{(\text{log IC}_50 - \text{HillSlope})}} \]  

(17)

where y corresponds to $\Theta_9$-bound, $y_{\max}$ and $y_{\min}$ represent the lower and the upper asymptotic value of the sigmoidal function, respectively. x corresponds to log [9]₀, log IC₅₀ represents the logarithm of the IC₅₀ concentration, and HillSlope determines the steepness of the curve. We set $y_{\min}$ to 0, because we assumed that at large concentrations of 9, all CA II molecules would be bound to 9 and hence $\Theta_9^{\text{bound}}$ would be 0. In order to obtain the value of y_{max}, we calculated $\Theta_9^{\text{bound}}$ before addition of 9 (i.e., [9]₀ = 0) in two steps. We first used eq 13 to calculate [8]_{eq} by employing the previously determined value of K₉₉ = 6.7×10⁻³ M⁻¹ and a value of [CA II]₀ = 9.3×10⁻⁹ as well as a value of [8]₀ = 16.1 nM; with these values we obtained [8]_{eq} = 2.2×10⁻⁸ M. Second, by using this value for [8]_{eq} and eq 16, we obtained a value of K₉₉ = 0.86. Consequently, we used this value of 0.86 for y_{max} to perform the best curve fit of the blue points in Figure 7 with eq 17. This analysis generated the following two values: log IC₅₀ = −4.89 ± 0.04, and HillSlope = 2.22 ± 0.44 (R² = 0.981, N = 4). Hence, the mean value for the IC₅₀ concentration for displacement of 8 from CA II by adding 9 was 12.9 µM.

Second, from this value of IC₅₀ we calculated the concentration of 9 that was free in the equilibrium mixture [9]_{eq} when the total concentration of 9, [9]₀, was equal to the IC₅₀ (i.e., [9]₀ = 12.9 µM) with the following equation:

\[ [9]_{\text{eq}} = \frac{\text{IC}_50 - [\text{CA II}]_0}{2} \left( \frac{[8]_{\text{eq}}}{K_{\text{d8}}} + \frac{[\text{CA II}]_0}{K_{\text{d8}} + [8]_{\text{eq}} + [\text{CA II}]_0} \right) \]  

(18)

For this calculation, we first computed the value of [8]_{eq} when [9]₀ = 12.9 µM in two steps: (i) eq 14 yielded a value of Q₃₀ = 5.7% when [9]₀ = 12.9 µM and (ii) from this value of Q₃₀, eq 7 yielded the value of [8]_{eq} = 8.2 nM. The remaining two values that we used to calculate [9]_{eq} with eq 18 were K₉₉ = 1.5 µM and [CA II]₀ = 9.3 µM. With these parameters, eq 18 returned a value of [9]_{eq} = 4.8 µM.

And in the third and final step, we used the value of [9]_{eq} = 4.8 µM and the following formula derived by Linden to calculate the desired value of K₉₉:

\[ K_{\text{d9}} = \frac{[9]_{\text{eq}}}{1 + \frac{[8]_{\text{eq}}}{K_{\text{d8}}}} \left( \frac{[\text{CA II}]_0}{K_{\text{d8}} + [8]_{\text{eq}} + [\text{CA II}]_0} \right) \]  

(19)

with [8]_{eq} = 8.2 nM, K₉₉ = 1.5 µM, and [CA II]₀ = 9.3 µM, we obtained a value for K₉₉ = 0.7 µM. If we repeated this analysis by employing the range of values for K₉₉ that we determined previously (K₉₉ = 0.9 – 2.8 µM), we obtained a range from 0.4 to 1.2 µM for the dissociation constant K₉₉. The value of 0.7 µM for K₉₉ is in excellent agreement with the dissociation constant of 0.73 ± 0.02 µM determined for the same interaction (i.e., for binding of 4-carboxybenzenesulfonamide (9) to CA II) by isothermal titration calorimetry.

Practicability of Quantifying Protein–Ligand Interactions with Ion Channel-Forming Peptides. In asking if the approach described here offers a broadly applicable assay platform, we see three major challenges. First, the preparation of planar bilayers of high quality requires expertise, experimental skill, and appropriate equipment, which includes a low noise current amplifier, a data acquisition board with adequate software, a Faraday cage, and typically a vibration-damping platform. Second, planar lipid bilayers are inherently metastable systems; their lifetime typically ranges from a few minutes to a few hours. When membranes broke, it was often necessary to wash the
entire setup and start a new experiment. Because the analysis presented here was based on equilibrium conditions of binding, it took several hours to record entire dose–response curves. Hence, in some cases only a part of the dose–response curves could be completed with a given bilayer. Third, it can be difficult to keep the bilayer area of a planar lipid membrane constant over minutes to hours. Fluctuations in bilayer area (due to fluctuations of the area and thickness of the torus) could cause fluctuations in transported charge, and these fluctuations could occur within one dose–response curve, potentially leading to inaccuracies of \( Q \). For each new bilayer, we measured the capacitance of the membrane, a quantity that is proportional to the bilayer area, and waited until the capacitance appeared to reach a nearly constant value. For some membranes, this waiting period could take tens of minutes. Then we measured \( Q_{\text{max}} \) for this particular membrane before adding CA II or 9. In some cases, we measured only a limited number of concentrations of a dose–response curve and then purposely prepared a new membrane to minimize possible fluctuations of the membrane area over long periods of time.

The assays presented here, although promising and remarkably accurate with respect to the determined dissociation constants, are thus, in their current configuration, not practical for real-world sensor applications; they are best performed under well-defined conditions in research laboratories relatively proficient in ion-channel biochemistry. One potential strategy to reduce problems with changing bilayer areas might be to perform the assays on apertures with large diameters (e.g., 500 \( \mu \)m) rather than the small diameters that we chose (<50 \( \mu \)m). If these large apertures would be prepared in very thin polymer films, then the surface area of the membrane that would be occupied by the torus would be relatively small compared to the bilayer area over these large pores. Fluctuations in the area of the torus may thus have a relatively small effect. Planar lipid bilayers over large diameter pores are, however, less stable than bilayers over small pores (which is the reason why we chose small pores in this work), and the large capacitance of these membranes generates large current noise in the recordings. Recent developments of hydrogel-embedded planar lipid bilayers may, however, make it possible to extend the lifetime of these bilayers.

**Conclusion**

Using a system of CA II and alamethicin covalently attached to a sulfonamide ligand, we demonstrated that binding of a protein to an ion channel-forming peptide that carries a ligand can be used to quantify protein–ligand interactions by measuring the inhibition of ion channel activity upon binding. Time-averaging of the current transported through the bilayer made it possible to determine the dissociation constant for this monovalent protein–ligand interaction. The resulting assay could be carried out in small volumes of solution (here 3 mL but volumes smaller than 50 \( \mu \)L have been used for planar lipid bilayer recordings of alamethicin activity) and required nanomolar concentrations of the ion channel-forming peptide (here alamethicin sulfonamide) as well as low micromolar concentrations of the protein (here CA II).

To make this type of ion channel-based affinity sensors practical, the synthetic route for attaching ligands to alamethicin will have to be simplified, ideally to a degree that it could be carried out by nonchemists. The work presented here is only a first step in this direction because it starts from commercially available building blocks. The overall yield was only moderate, and the purification required expertise in preparative HPLC techniques. Further requirements are that the ligands should have low molecular weight and must be chemically amenable to covalent coupling to the pore-forming peptide; they also must be sufficiently stable to withstand the coupling reaction without changing their physicochemical properties. As with most applied experiments on planar lipid bilayers, the mechanical and thermodynamic stability of the bilayers would have to be improved before such sensors will yield widely used, robust, and user-friendly assays. Quantitative analyses of changes in the transported charge through pores of ion channel-forming peptides as a function of the addition of an analyte pose an additional challenge: the area of the lipid bilayer has to be kept as constant as possible or correction strategies have to be developed that take into account possible fluctuations in bilayer area over time. One possible strategy may be frequent measurements of the electrical capacitance of the bilayer and normalizing the relative transported charge over the membrane capacitance (or the bilayer area, which is proportional to the membrane capacitance). Another strategy may be to embed the bilayer into a hydrogel polymerized in situ. The hydrogel can stabilize the torus of the membrane and thus minimize fluctuations in area of the bilayer.

We believe that the active research efforts in the area of microfabricated planar lipid bilayer setups, as well as in the area of synthesizing functional derivatives of ion channel-forming peptides, will help to overcome, at least to some extent, these challenges of affinity sensors based on ion channels. If so, these sensors might make it possible to perform quantitative affinity assays in volumes below 50 \( \mu \)L and at concentrations of ligand and receptor below 10 \( \mu \)M. Their advantage might be to require only subnanomole amounts of receptor and ligand.

In comparison to existing methods such as isothermal titration calorimetry, the method presented here has the advantage that it is a single molecule method in the sense that the flux of ions through a single assembled pore can be monitored readily and with a time resolution below milliseconds. Another advantage

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(102) Fertig, N.; Blick, R. H.; Behrends, J. C. Biophys. J. 2002, 82, 3056–3062.


compared to isothermal titration calorimetry is that sensing with ion channels requires significantly lower concentrations of ligand (here 8) and the detection mechanism is, in principle, strikingly simple (i.e., applying a small constant voltage and measuring a small current over time). In addition, the footprint of the entire experimental setup can be small making this method attractive for potential applications as portable sensors.113–115

Moreover, sensing with ion channels is an amplifying method and has some analogy to standard affinity assays such as enzyme-linked immunosorbent assays (ELISAs), which utilize the amplification provided by enzymatic turnover. This method takes advantage of the inherent amplification of ion channels in the sense that one channel can conduct thousands of ions per millisecond. Another advantage of channel-based sensing is that it is not sensitive to colored samples or samples that might contain quenchers of fluorescence or inhibitors of the enzymes that are used for amplification.63 In addition, ion channel-based sensing can, presumably, proceed in solution as long as the affinity interaction reduces the concentration of free channel-forming peptide and thus shifts the equilibria in Figure 1 in the direction of reduced concentrations of peptide in a functional pore. ELISA assays, in contrast, are inherently surface-based assays.

Nonetheless, at this stage, ELISA assays provide superior robustness and ease of use compared to the channel-based affinity assay demonstrated in this report. Compared to isothermal titration calorimetry, sensing with ion channels has the disadvantage that it requires derivatization of the ion channel-forming peptide to attach each ligand of interest. Moreover, as opposed to isothermal titration calorimetry, sensing based on ion channels does not provide thermodynamic quantities such as changes in enthalpy. Compared to both methods, ELISA and isothermal titration calorimetry, sensing with ion channels, at this stage of development, is hampered by the stability of the bilayers, which is limited to periods of minutes to hours.

To carry out the analysis presented here, we proposed that the disruption and restoration of self-assembly of a derivative of alamethicin can be used for quantifying receptor ligand interactions. Although we did not provide definite proof for this mechanism of detection, we think that the excellent agreement of the determined dissociation constants with literature values support this mechanism. On the basis of these results, we suggest that utilizing the disruption or restoration of self-assembly for sensing applications is a fundamentally interesting and promising avenue because self-assembly is often a cooperative process and may thus respond strongly to specifically engineered disturbances.

Experimental Section

Materials. All chemicals, including bovine CA II (CA II, p 5.9, E.C. 4.2.1.1), were purchased from Sigma-Aldrich (St. Louis, MO) unless stated otherwise. Mass spectra were performed by matrix-assisted laser desorption/ionization mass spectrometry (MALDI-TOF) using α-cyano-4-hydroxy-cinnamic acid as matrix. All analytical HPLC separations were run with a Microsorb C18 column 5 µm (4.6 × 250 mm) using a linear gradient from A (100% water containing 0.1% TFA) to B (100% acetonitrile containing 0.08% TFA), at a flow rate of 1.2 mL min⁻¹ (UV detection at 214 and 254 nm). Alamethicin with a purity >90% (HPLC) was purchased from Sigma (St. Louis, MO). We started with commercial alamethicin (6) (isolated from Trichoderma viride) and separated the two alamethicin variants (F30 and F50), by HPLC on a 5 µm C18 column (10 × 250 mm) using a linear gradient of 70% A with 30% B to 100% B over 40 min at a flow rate of 5 mL min⁻¹. We used the F30 fraction for all experiments; fractions containing alamethicin F30 were combined and evaporated to dryness (purity >99%). MS (MALDI-TOF) m/z: 1965.77 [M + H]+, calculated for C₂₇H₂₆N₇O₁₄S⁺ [M + H]+ 1965.32.

F30: Ac-Aib-Pro-Aib-Ala-Ala-Gln-Ala-Val-Ala-Gly-Leu-Aib-Pro-Ala-Aib-Glu-Gln-Phenylnalinalin

F50: Ac-Aib-Pro-Aib-Ala-Ala-Gln-Ala-Val-Ala-Gly-Leu-Aib-Pro-Ala-Aib-Glu-Gln-Phenylnalinalin

Compound 1. Acid 1 was prepared as described by Avila et al.116 The yield was 89%, and the product was pure. 1H NMR (400 MHz, DMSO-d₆) δ 1.53 (m, 4H), 2.17 (t, 2H), 2.32 (t, 2H), 3.58 (s, 3H), 4.32 (d, 2H), 7.33 (s, 2H), 7.41 (d, 2H), 7.77 (d, 2H), 8.43 (t, 1H), HRMS (FAB) m/z: 329.1181 [M + H]+, calculated for C₁₉H₁₉N₅O₄S⁺ [M + H]+ 329.1171.

Compound 2. Compound 2 was synthesized according to Roy and Mallik.117 1H NMR (400 MHz, MeOH/CDCl₃) δ 2.82 (t, 2H), 3.35–3.39 (m, 2H), 3.53–3.66 (m, 6H), 5.07 (s, 2H), 7.31–7.34 (m, 5H).

Compound 3. To a solution of (Boc-aminooxy)acetic acid (0.437 g, 2.29 mmol), N-ethyl-N′-dimethylaminopropylcarbodiimide.HCl (0.438 g, 2.29 mmol) and N,N-diisopropylethylamine (1.09 mL, 6.24 mmol) in dimethylformamide (5 mL) was added 2 (0.587 g, 2.08 mmol), and the solution was stirred at room temperature for 5 h. The solution was evaporated, and the product partitioned between ethyl acetate and water. The organic layer was washed with water (2 × 50 mL), dried over sodium sulfate, and evaporated to dryness. The crude compound was chromatographed (SiO₂, using 100% ethyl acetate as eluent) to yield 3 (350 mg, 1.16 mmol, 56% yield) as a clear oil. 1H NMR (500 MHz, CDCl₃) δ 1.46 (s, 9H), 1.57–1.69 (m, 4H), 3.29 (t, 2H), 3.43 (t, 2H), 3.54 (t, 2H), 3.57 (t, 2H), 3.61 (s, 4H), 4.25 (s, 2H), 5.07 (s, 2H), 7.25–7.38 (m, 5H); MS (MALDI-TOF) m/z: 478.74 [M + Na]+, calculated for C₂₇H₂₆N₅O₄Na⁺ [M + Na]+ 478.51.

Compound 4. Compound 4 (157 mg, 0.34 mmol) and 10% Pd/C (20 mg) were combined in ethanol (20 mL). This mixture was hydrogenated at atmospheric pressure and room temp for 2 h. The mixture was then filtered through Celite, the filter cake was washed with 2 × 10 mL of ethanol, and the resulting solution was concentrated by evaporation. The acid 1 (119 mg, 0.38 mmol), N-ethyl-N′-dimethylaminopropylcarbodiimide.HCl (73 mg, 0.38 mmol) and N,N-diisopropylethylamine (180 mL, 1.03 mmol) were combined in 5 mL of DMF at room temperature with stirring, and the reaction was allowed to proceed at room temperature overnight. The crude product was purified by HPLC (linear gradient, 100% A to 80% B, over 40 min) and lyophilized to afford 4 (111 mg, 0.18 mmol, 52% yield) as a clear oil. 1H NMR (500 MHz, CDCl₃) δ 1.47 (s, 9H), 1.57–1.69 (m, 4H), 2.22 (t, 2H) 2.28 (t, 2H), 3.35 (t, 2H), 3.44 (t, 2H), 3.54 (t, 2H), 3.62 (s, 4H), 4.25 (s, 2H), 4.3 (s, 2H), 4.74 (d, 2H), 7.42 (d, 2H), 7.85 (d, 2H); analytical HPLC tR 12.77 min (linear gradient, 100% A to 100% B, over 20 min); MS (MALDI-TOF) m/z: 618.27 [M + H]+, calculated for C₃₀H₂₄N₅O₁₂S⁺ [M + H]+ 618.72.

Alamethicin-Sulfonamide, 8. To a solution of alamethicin F30 (8 mg, 4.08 µmol) in dichloromethane, Dess-Martin periodinane (17.3 mg, 40.82 µmol) was added.118 The mixture was filtered and concentrated to a solid by evaporation. The crude peptide derivative was purified by preparative HPLC (linear gradient, 70% A with 30% B to 100% B, over 40 min), and the solution evaporated to afford alamethicin aldehyde 7 (4 mg, 2.04 µmol) as a white solid. No impurities were detected by

analytical HPLC (linear gradient, 100% A to 100% B, over 20 min). The Boc-protected compound 4 (6.3 mg, 10.21 μmol) was dissolved in 10 mL of trifluoroacetic acid and stirred for 30 min at room temperature. The solution was evaporated to dryness. The residue was dissolved in methanol (1 mL) and 7 (4 mg, 2.04 μmol) was added. The reaction was allowed to proceed for 2 h and then evaporated. The crude peptide derivative was purified by HPLC (linear gradient, 70% A with 30% B to 100% B, over 40 min) and evaporated to afford 8 (2.4 mg, 0.97 μmol, 24% yield) as a white solid. HPLC t_R = 14.20 and 14.42 min (linear gradient, 70% A with 30% B to 100% B, 20 min); MS (MALDI-TOF) m/z 2484.38 [M + Na]^+ , calculated for C_{113}H_{181}N_{27}NaO_{32}S^+ [M + Na]^+ 2484.86.

**Molecular Modeling.** We generated the energy minimized structure shown in Figure 3B and C using molecular mechanics calculations employing AMBER force field parameters in water (MacroModel software, version 7.5, Schroedinger Inc.). We constructed employing AMBER force field parameters in water (MacroModel shown in Figure 3B and C using molecular mechanics calculations

5 residues of the alamethicin peptide (from the Protein Data Bank) in MacroModel. For this calculation, the

6 by modification of the C-terminus of the crystal structure of 6 (1AMT from the Protein Data Bank) in MacroModel. For this calculation, the residues of the alamethicin peptide (6) were fixed during the conformational analysis. We allowed the ligand (5) to rotate freely during these calculations. After performing 5000 iterations of conformational analysis of 8, we selected the lowest energy conformation.

**Planar lipid Bilayer Experiments.** We used the same experimental setup and procedure as described in detail by Mayer et al.49 Briefly, we prepared Teflon AF films by melting a solution of 6% Teflon AF in Fluorinert FC-75 solvent (DuPont Fluoroproducts, Wilmington, DE) around a sharp tip (gold-plated tungsten probe tips with a nominal tip diameter of 10 μm from Lucas Signatone, Gliroy, CA) that was oriented perpendicular to, and placed in contact with, a silicon wafer such that the tip rested on the wafer by gravity. Before casting the Teflon AF solution, the wafer was treated with an air plasma followed by silanization with tridecafluoro-1,1,2,2-tetrahydrooctyl-1-trichlorosilane (United Chemical Technologies, Bristol, PA) under vacuum (60 mmHg) to facilitate the removal of the film of Teflon AF from the wafer after evaporation of the solvent.

Before using these Teflon AF sheets for planar lipid bilayer experiments, we treated10 the area close to the pores with a solution of 5% squalene (Sigma, St. Louis, MO) in pentane (Sigma-Aldrich, St. Louis, MO) by dipping the tip of a tissue paper first into the squalene solution and then moving it over the area close to the pore in the Teflon film.10 We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF support. The result of this procedure is a lipid bilayer that their acyl chains face each other and span over the aperture in the Teflon AF film. We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF film. We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF film.

To facilitate the removal of the film of Teflon AF from the wafer after casting the Teflon AF solution, the wafer was treated with an air plasma followed by silanization with tridecafluoro-1,1,2,2-tetrahydrooctyl-1-trichlorosilane (United Chemical Technologies, Bristol, PA) under vacuum (60 mmHg) to facilitate the removal of the film of Teflon AF from the wafer after evaporation of the solvent.

We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF film. We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF film. We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF film. We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF film. We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF film. We formed all bilayers by the “folding technique” in which two monolayers of lipids at an air-electrolyte interface are raised such that their acyl chains face each other and span over the aperture in the Teflon AF film.