Criterion for Amino Acid Composition of Defensins and Antimicrobial Peptides Based on Geometry of Membrane Destabilization

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Supporting Information

ABSTRACT: Defensins comprise a potent class of membrane disruptive antimicrobial peptides (AMPs) with well-characterized broad spectrum and selective microbicidal effects. By using high-resolution synchrotron small-angle X-ray scattering to investigate interactions between heterogeneous membranes and members of the defensin subfamilies, α-defensins (Crp-4), β-defensins (HBD-2, HBD-3), and θ-defensins (RTD-1, BTD-7), we show how these peptides all permeabilize model bacterial membranes but not model eukaryotic membranes: defensins selectively generate saddle-splay (“negative Gaussian”) membrane curvature in model membranes rich in negative curvature lipids such as those with phosphoethanolamine (PE) headgroups. These results are shown to be consistent with vesicle leakage assays. A mechanism of action based on saddle-splay membrane curvature generation is broadly enabling, because it is a necessary condition for processes such as pore formation, blebbing, budding, and vesicularization, all of which destabilize the barrier function of cell membranes. Importantly, saddle-splay membrane curvature generation places constraints on the amino acid composition of membrane disruptive peptides. For example, we show that the requirement for generating saddle-splay curvature implies that a decrease in arginine content in an AMP can be offset by an increase in both lysine and hydrophobic content. This “design rule” is consistent with the amino acid compositions of 1080 known cationic AMPs.

1. INTRODUCTION

Antimicrobial peptides (AMPs) comprise an important component of the innate host defense system. Collectively, AMPs have broad spectrum antimicrobial activity. Although AMPs are diverse in sequence and structure, most share two general structural features: they are amphipathic and cationic. From in vitro studies, AMP amphipathicity is inferred to disrupt membranes through a combination of electrostatic interactions of the cationic AMP with the anionic membrane followed by insertion of hydrophobic patches into the nonpolar interior of the bilayer. Membranes can be disrupted by AMPs via a diverse range of processes, including pore formation, blebbing, budding, and vesicularization. This diversity of outcomes has impeded the development of a detailed, molecular understanding of how AMPs destabilize bacterial membranes specifically. Such knowledge can in principle provide “design rules” that impact development of synthetic antimicrobials.

Mammals produce two main types of membrane active AMPs: defensins and cathelicidins. Defensins are synthesized by varied epithelia and by phagocytes and function as a biochemical line of defense against microbial infection. Vertebrate defensins share general features, including cationic charge, MW of 2–5 kDa, distinctive trisulfide arrays, and β-strand structures constrained by the disulfide pairings. They can be separated into three subfamilies, α-defensins, β-defensins, and θ-defensins. Because many aspects of defensin biology are known, this is an ideal prototypical family of AMPs to investigate the detailed molecular mechanisms responsible for selective activity against bacterial but not mammalian membranes.

In this work, we use synchrotron small-angle X-ray scattering (SAXS) to map out the fundamental curvature deformation modes induced in model cell membranes by defensins and elucidate causative relationships between curvature deformations and membrane disruption processes. Mammalian defensins are systematically investigated, using members from the

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three defensin subfamilies, including rhesus monkey \( \theta \)-defensin-1 (RTD-1), baboon \( \theta \)-defensin-7 (BTD-7), mouse Paneth cell \( \alpha \)-defensin cryptdin-4 (Crp-4), human \( \beta \)-defensin-2 (HBD-2), and human \( \beta \)-defensin-3 (HBD-3). The bactericidal activity of these defensins is correlated with the induction of saddle-splay (equivalently negative Gaussian) membrane curvature, which enables membrane destabilizing processes.\(^{13}\) All peptides restructured vesicles into porous bicontinuous phases rich in saddle-splay curvature when lipid compositions mimic those of bacterial membranes, but not when the lipid compositions are more representative of mammalian membranes. A key parameter for activity is the concentration of negative intrinsic curvature \( (\theta_0 < 0) \) lipids, such as those with PE headgroups, which exist at significantly higher concentrations in bacterial cytoplasmic membranes as compared to eukaryotic membranes. The existence of homologous behavior in synthetic antimicrobials\(^{14–16}\) suggests a common root mechanism for selective membrane permeation. In general, AMPs destabilize membranes via a diverse range of processes, many of which involve saddle-splay curvature. We deduce a criterion for amino acid compositions of AMPs based on the requirement for generating saddle-splay membrane curvature and show that it is consistent with trends in amino acid composition of 1080 known cationic AMPs.

2. EXPERIMENTAL SECTION

2.1. Preparation of Peptides. Recombinant Crp4 was expressed and purified as His6-tagged fusion proteins as described.\(^{17–19}\) Expression of recombinant fusion proteins was induced by adjusting exponentially growing \( E. \) coli BL21-CodonPlus (DE3)-RIL cells to 0.1 mM isopropyl-\( \beta \)-D-thiogalactopyranoside and incubating at 37 °C for 6 h in Terrific Broth as described in earlier reports. Cells were lysed by sonication in 6 M guanidine-HCl, 100 mM Tris (pH 8.0), and the suspension was clarified by centrifugation. His6-tagged fusion proteins purified by nickel-nitrilotriacetic acid resin affinity chromatography (Qiagen, Valencia, CA) were cleaved with cyanogen bromide and purified by analytical C18 reverse phase high pressure liquid chromatography (RP-HPLC). Peptide homogeneity was confirmed in analytical acid-urea polyacrylamide gel electrophoresis (AU-PAGE), a highly sensitive measure of defensin foldamers. Molecular masses were verified by matrix-assisted laser desorption ionization-time-of-flight mass spectrometry (MALDI-TOF MS), and peptides were quantified using extinction coefficient calculations at 280 nm performed at ExPASY.\(^{20}\) \( \theta \)-Defensins RTD-1 and BTD-7 were produced by solid-phase synthesis, purified, and characterized as described previously.\(^{21,22}\) HBD-2 and HBD-3 are expressed and purified as described in refs 23,24.

2.2. Liposome Preparation for X-ray Measurements. DOPC (1,2-dioleoyl-sn-glycero-3-phosphocholine), DOPE (1,2-dioleoyl-sn-glycero-3-phosphoethanolamine), DOPG \( \{1,2\)-dioleoyl-sn-glycero-3-phospho-rac-(1-glycerol) \} (sodium salt), and DOPS \( \{1,2\)-dioleoyl-sn-glycero-3-phospho-1-serine \} (sodium salt), lyophilized lipids from Avanti Polar lipids, were used without further purification. For X-ray experiments, we prepared small unilamellar vesicles (SUVs) by sonication. DOPG, DOPS, DOPE were dissolved in chloroform/methanol, and DOPC, DOPE were dissolved in chloroform for stock solutions. Mixture of DOPG and DOPE were prepared at 1:4 mass ratio to approximate bacterial lipid compositions, while DOPS, DOPE, and DOPC at 1:4:0 and 1:2:2 mass ratios were used to approximate the membranes of eukaryotic cells. Lipid mixtures were dried under \( N_2 \), desiccated overnight under vacuum, and resuspended at a final concentration of 20 mg/mL in Millipore H\(_2\)O. Solutions were incubated at 37 °C overnight and then sonicated until clear. SUVs were obtained via extrusion (0.2 μm pore Nucleopore filter).

2.3. SAXS Experiments. Defensin and lipid solutions were thoroughly mixed at specific peptide to lipid ratios (P/L) and sealed in quartz capillaries. All samples were prepared in 100–150 mM NaCl to simulate physiological salt conditions. For SAXS experiments at the Stanford Synchrotron Radiation Laboratory (BL4- 2) and at the Advanced Photon Source (BESSRC-CAT BL-12IDC), monochromatic X-rays with energies of 9–11 and 12 keV, respectively, were used. Scattered radiation was collected using a MAR Research CCD area detector (pixel size, 79 μm). Samples were checked for consistency by multiple measurements with different X-ray sources. No radiation damage was observed for the incident beam intensities and the exposure times used. 2D SAXS powder patterns were integrated using the Nika 1.2 package\(^{25}\) and FIT2D.\(^{26}\)

2.4. Vesicle Dye Leakage Experiments. Giant unilamellar vesicles (GUVs) were prepared using the swelling method. Stock solutions of lipids DOPS, DOPE, and DOPC in chloroform were deposited (100 μL at 20 mg/mL) at the desired mass ratios onto roughened, cleaned Teflon, along with dye DiO (10 μL at 1 mg/mL). After drying under vacuum, the Teflon was hydrated with moist nitrogen gas for 30 min, and then 5 μL of swelling solution was added (100 mM sucrose plus 20 μM Alexa 633 maleimide) and incubated for 2 days at 37 °C. After swelling, the GUV suspension was diluted 40× into 200 mM glucose. 200 μL of this dilution was placed on a coverslip soaked in 1% BSA, and Crp4 (10 μL at 1 mg/mL) was added. A Leica SP2 laser scanning confocal microscope was used.

2.5. Analysis of Data from the Antimicrobial Peptide Database.\(^{27}\) Data was obtained via the online antimicrobial peptide database.\(^{28}\) Comparison of Helical Peptides with Defensins. Eligible peptides were required to be active against Gram-negative or Gram-positive bacteria. A pool of 299 peptides with helix structure is compared to a pool of 143 defensins. “Average % of AMP composition” is the number of times an amino acid is found in a pool divided by the total number of amino acids in that pool.

Lysine/Arginine Ratio versus % Hydrophobicity. Eligible peptides were required to have a net positive charge and be active against either Gram-negative or Gram-positive bacteria. Percentage of lysine, %K, and percentage of arginine, %R, were read off from the sequence information for a set hydrophobic bin. Here, %K (%R) = number of lysines (arginines) found in the bin/total number of amino acids in the bin. Note that %K/%R = N\(_K\)/N\(_R\) (defined below).

Lysine/Arginine Ratio versus (Hydrophobicity). Eligible peptides were required to have a net positive charge and be active against either Gram-negative or Gram-positive bacteria. These criteria retrieved a pool of 1080 peptides. For a given peptide, \( j \), its average hydrophobicity is defined by:

\[
\text{(Hydrophobicity)}_j \equiv \frac{1}{n} \sum_{i=1}^{n} w_i
\]

where \( n \) = number of amino acids in the peptide, and \( w_i \) = the hydrophobicity of the \( i \)th amino acid in the peptide. This value is set by the particular hydrophobicity scale used. To facilitate easier comparison across scales, the value of the apparent free energy of membrane insertion for an amino acid, \( \Delta G_{\text{app}}^{\text{mem}} \), based upon the Willmer–White biological scale, was reversed in sign, that is, \( \Delta G_{\text{app}}^{\text{mem}} \rightarrow -\Delta G_{\text{app}}^{\text{mem}} \). The published hydrophobicity values were used for all other scales.

To partition each peptide into a bin, the hydrophobic extremes for the pool were determined and used to set the range of hydrophobicity scale. The range was divided into 100 equal bins. For the M peptides in
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3. RESULTS

3.1. \( \theta \)-Defensins Generate Saddle-Splay Curvature in Model Bacterial Membranes with High PE Concentrations but Not in Model Eukaryotic Membranes with Low PE Concentrations. \( \theta \)-Defensins are short 18 amino acid circularized peptides expressed only in old world monkeys and orangutans.\(^{21,29}\) Previous studies have shown that rhesus \( \theta \)-defensin-1 (RTD-1) and baboon \( \theta \)-defensin-7 (BTD-7) have activity against Gram-positive and Gram-negative bacteria and fungi.\(^{21,29}\) Moreover, RTD-1 has antiviral activity against HIV-1\(^{30}\) and the herpes simplex virus.\(^{31}\) Both \( \theta \)-defensins are amphi- pathic and contain five arginine residues.

Small unilamellar vesicles (SUbs) with phospholipid compositions characteristic of Gram-negative bacteria (DOPG/DOPC = 20/80) were incubated with specific defensins to study the induced structural changes. Synchrotron SAXS profiles from lipid solutions only showed a broad characteristic feature consistent with a single lipid bilayer form factor expected for unilamellar vesicles (see Supporting Information, Figure S1). When exposed to RTD-1 or BTD-7 (Figure 1A,B), the vesicles underwent a drastic structural transition and displayed characteristic correlation peaks with ratios \( \sqrt{2:3:4:6} \), which indicate the formation of a cubic \( Pn3m \) “double-diamond” lattice. By fitting the slopes of the measured Q-values at every peak, we calculate lattice parameters, \( d_{RTD-1} = 16.00 \text{ nm} \) and \( d_{BTD-7} = 15.98 \text{ nm} \), for BTD-7 and RTD-1, respectively.\(^{32}\) The \( Pn3m \) (Figure 1C) is a bicontinuous cubic phase where two nonintersecting water channels are separated by a lipid bilayer.\(^{33}\) The center of the bilayer traces out a minimal surface with negative Gaussian curvature at every point. Although bilayer saddle-splay curvature is different from the self-connected monolayer saddle-splay curvature seen in a toroidal pore (Figure 1D), it should be noted that each constituent monolayer in a bicontinuous phase also possesses negative Gaussian curvature at every point and is equally accessible by AMPs. On such saddle-shaped surfaces (Figure 1C), the surface curves upward in one direction and downward in the perpendicular direction. For example, saddle-shaped surfaces are found inside toroids, which have a single hole each. This type of curvature is not found on the surfaces of spheres, which lack holes. Furthermore, protrusions such as buds and blebs (Figure 1D) require bilayer saddle-splay curvature at their base. Saddle-splay curvature is required for membrane poration, as well as for formation of different AMP-induced destabilization mechanisms depicted in Figure 1D.

To examine the role of membrane intrinsic curvature on induced saddle-splay curvature, we constructed ternary lipid membranes of constant anionic charge (DOPG = 20%) but with varying ratios of DOPC (\( \theta_c < 0 \)) and DOPE (\( \theta_c \approx 0 \)). In contrast to effects on PE-rich membranes, RTD-1 and BTD-7 did not induce structures that mimic eukaryotic membranes with low concentrations of PE. Phases rich in saddle-splay curvature were not induced by RTD-1 or BTD-7 in membranes with DOPG: DOPE:DOPC = 20:40:40 (Figure 1A,B), as indicated by the absence of their characteristic correlation peaks. The diffraction data showed a combination of broad form factor scattering characteristic of isolated bilayers and a lamellar phase with a period of \( d_{RTD-1} = 5.67 \text{ nm} \) and \( d_{BTD-7} = 5.67 \text{ nm} \) for RTD-1 and BTD-7, indicating that weak intermembrane attraction is induced, although curvature is not.

It is important to discriminate between necessary and sufficient conditions for saddle-splay curvature generation. Alamethicin,\(^{34}\) gramicidin S,\(^{35}\) and protegrin-1\(^{36}\) form cubic phases in pure PE lipids under specific processing conditions such as temperature cycling. The above results on defensins allow us to assess systematically the roles played by different lipid species. It has been previously proposed that anionic lipids in bacterial membranes are important for AMP function. We agree with this assessment. However, while anionic PG lipids and high concentration PE lipids in the target membrane are both necessary conditions for induction of saddle-splay curvature, neither by itself is a sufficient condition (see Supporting Information, Figure S1).
Figure 2. The α-defensin Crp-4 showed phase behavior similar to that of 3.2. Arginine-Rich α-Defensins Also Generate Saddle-Splay Curvature in Membranes Enriched with PE and Selectively Permeate PE-Rich GUVs. To investigate whether other arginine-rich defensins exhibit similar behavior, we conducted similar experiments on mouse cryptdin-4 (Crp-4), a 32 amino acid α-defensin expressed in mouse Paneth cells with in vitro microbicidal activity against Gram-negative and Gram-positive bacteria. Crp-4 is amphipathic, and 7 of the 10 cationic amino acids in Crp-4 are arginine residues.

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Like the α-defensins, the α-defensin Crp-4 generates saddle-splay curvature in model bacterial membranes with high PE but not in model eukaryotic membranes with low PE. Incubation of Crp-4 with SUVs of high PE lipid composition produced a Pn3m cubic phase with lattice parameter, a = 11.1 nm (Figure 2A), while reduction of membrane DOPE content led to the disappearance of the Pn3m cubic phase. Interestingly, previous work suggests that negative intrinsic curvature lipids can promote pore formation through modification of the Gaussian modulus. Typically the Gaussian curvature modulus is negative, which exacts a free energy penalty for pore formation. However, the presence of c0 < 0 lipids (such as PE lipids) changes the bilayer Gaussian curvature modulus toward positive values. The decreased free energy barrier to changes in membrane topology toward porous phases is consistent with our observation that bacterial membranes rich in negative intrinsic curvature lipids are more susceptible to pore formation.

Our X-ray experiments measure AMP-induced curvature deformations in a 3D lipid system, which has the freedom to connect into bulk phases such as the Pn3m. Both eukaryotic and prokaryotic cells have quasi-2D membranes, where induced saddle-splay curvature deformations can take the form of pores, blebs, or buds. To determine how peptide-induced saddle-splay curvature is manifested in 2D membranes, we investigated the effect of Crp-4 on giant unilamellar vesicles (GUV) by confocal microscopy. Liposomes with lipid compositions mimicking the membranes of bacteria (PS/PE = 20/80) and of eukaryotic cells (PS/PC = 20/80) were loaded with maleimide dye and incubated with Crp-4 (molar ratio P/L = 1; see Experimental Section). Dye leakage was evident in PE-rich membranes after peptide exposure (Figure 2B), consistent with Crp-4-induced pore formation in the intact vesicles. In contrast, negligible dye leakage was observed in PC-rich vesicles (Figure 2C). Selective permeation of GUVs rich in PE but not PC is consistent with the observation of pore-enabling Crp-4-induced saddle-splay curvature (Figure 2A) from SAXS experiments. Taken together, these results indicate Crp-4 preferentially creates pores in membranes with negative intrinsic curvature lipids.

3.3. General Mechanism for Defensins and Relation of Curvature Generation to Antimicrobial Peptide Sequence: HBD-2 and HBD-3 as Examples. We hypothesized the selective induction of saddle-splay curvature in model bacterial membranes is a generic feature among defensins. We used human β-defensin-2 (HBD-2, DEFB4) and human β-defensin-3 (HBD-3, DEFB103A) to test this hypothesis. HBD-2 is a 41AA amphipathic peptide, which is predominantly effective against Gram-negative bacteria. HBD-3 is 45AA, amphipathic, and generally more potent than HBD-2. Unlike the other two defensin subfamilies, which strongly favor arginine over lysine, β-defensins have intermediate arginine to lysine ratios. In HBD-3, the cationic residues consist of 7 arginines and 6 lysines, and HBD-2 has 2 arginines, 5 lysines, and 1 histidine.

Figure 3 shows the SAXS spectra of model bacterial and model eukaryotic membranes after exposure to HBD-2 and HBD-3. The capacity for membrane-specific saddle-splay curvature generation is observed and is consistent with behavior of β-defensins and α-defensins. Both β-defensins showed weak interactions with membranes enriched with DOPC and produced lamellar phases in membranes with intermediate amounts of PE/PC (Figure 3A,B). In DOPS/DOPC = 20/80 membranes, HBD-2 generated a Pn3m cubic phase, with lattice parameter a = 13.8 nm (Figure 3A). HBD-3 also generated saddle-splay curvature in DOPS/DOPC = 20/80 membranes, which is exhibited in reflections with ratio, 6:8:14:16, characteristic of the cubic Ia3d “gyroid” phase with a = 18.0 nm (Figure 3B). Like the Pn3m, the Ia3d is a bicontinuous cubic phase consisting of two nonintersecting water channels separated by a single lipid bilayer, which has saddle-splay curvature everywhere on its minimal surface.

4. DISCUSSION

4.1. Average Induced Saddle-Splay Curvature, (K), from SAXS Measurements Allows Quantitative Comparison of Different Antimicrobial Peptides. Our previous work on cell penetrating peptides has shown lysine and arginine generate different types of membrane curvatures. Because lysine and arginine are cationic, they both induce wrapping by anionic membranes. This is due to the entropy of counterions condensed on both the charged membrane and the short charged polymers. Electrostatic compensation between a cationic polymer and an anionic membrane leads to the release of condensed counterions into the bulk solution and a corresponding large entropy gain, which results in strong polymer–membrane binding. This has been observed in a
The average Gaussian curvature to form cubic phases. Clearly more work, experimental, and computational, is required for a full understanding of these effects. Extant theoretical studies of saddle-splay curvature. In contrast, the amine group of arginine supports multidentate hydrogen bonding, which can organize bulky lipid head groups to generate positive curvature in the perpendicular direction, resulting in saddle-splay curvature. In contrast, the amine group of lysine can only hydrogen bond to one lipid headgroup at a time. Consistent with this, polylysine induces negative Gaussian curvature. Neither HBD-2 (A) or HBD-3 (B) produced cubic phases in membranes with low PE concentrations. In DOPS/DOPE/DOPC = 20/40/40 membranes, HBD-2 and HBD-3 restructured SUVs into lamellar phases with lattice spacings $d = 6.2$ and $7.0$ nm, respectively, and DOPS/DOPC = 20/80 membranes showed only weak interaction with both β-defensins. Conversely, in DOPS/DOPE = 20/80 membranes, HBD-2 induced coexisting $Pn3m$ and inverted hexagonal (lattice spacing $a = 7.5$ nm) phases, while HBD-3 induced an $Ia3d$ cubic phase.

The exchange of R with K and hydrophobicity is illustrated by comparing β-sheet defensins with α-helical AMPs. (A) On average, defensins contain much greater amounts of R than do α-helical AMPs and utilize less of each type of hydrophilic amino acid. (B) Overall, the proportion of hydrophilic amino acids is comparable in defensins and α-helical AMPs. The exception is K, which is more commonly found in α-helical AMPs. Blue denotes basic amino acids, while negatively charged amino acids are colored red.

Figure 3C shows the $\langle K \rangle$ values extracted from the diffraction data for arginine-rich defensins (RTD-1, BTD-7, Crp-4) and protegrin-1 (PG-1)60 (see Supporting Information, Figure S3). A correlation exists between negative Gaussian curvature and arginine content in structurally similar peptides. HBD-2 and HBD-3, which have ~50% lysines and ~50% arginines, clearly deviate from this curve, although $\langle K \rangle$ is more negative for HBD-3 as compared to HBD-2, consistent with the greater number of arginines in HBD-3.

4.2. The Requirement of Saddle-Splay Membrane Curvature Generation Places Constraints on Amino Acid Composition of Antimicrobial Peptides: The Saddle-Splay Curvature Selection Rule. Defensins can generate pores, but AMPs have been observed to disrupt membranes via processes that do not involve pores. Generation of the required saddle-splay membrane curvature for these processes places constraints on the amino acid composition and composite hydrophobic and cationic components of membrane disruptive peptides. Hydrophobic amino acids generate positive curvature. Nonpolar components of the amphipathic peptide are inserted into the membrane, which increases the hydrophobic volume of the perturbed lipid monolayer and thereby creates positive curvature strain. Differences in lipid headgroup chemistry will not drastically affect this mechanism.

Cationic amino acids work differently. We have demonstrated lipid headgroup organization by arginine and lysine generates distinct types of membrane curvature deformations. These

large number of systems. Because maximal contact between the peptide and membrane leads to maximal counterion release, the membrane will tend to wrap around the peptide generating negative curvature.53,54 The guanidine group of arginine supports multidentate hydrogen bonding, which can organize bulky lipid head groups to generate positive curvature in the perpendicular direction, resulting in saddle-splay curvature. In contrast, the amine group of lysine can only hydrogen bond to one lipid headgroup at a time. Consistent with this, polylysine induces negative mean curvature on membranes to form inverted hexagonal phases, while polyarginine can also induce saddle-splay curvature to form cubic phases. Clearly more work, experimental, theoretical, and computational, is required for a full understanding of these effects. Extant theoretical studies suggest the anisotropy associated with saddle-splay curvature deformations may also provide AMPs with a way to communicate and organize cooperatively. Currently, we are directly testing effects from peptide orientation and conformation, as well as proximity effects of different amino acids.

The $Pn3m$, $Ia3d$, and $Im3m$ cubic phases are related by a Bonnet transformation, so their average Gaussian curvatures can be quantitatively compared.55,56 The average Gaussian curvature is $\langle K \rangle = 2\pi\chi / a^2 A_{3c}$, where $a$ is the cubic lattice constant, $\chi$ is the Euler characteristic, and $A_{3c}$ is the surface area per unit cell. Figure 3C shows the $\langle K \rangle$ values extracted from the diffraction data for arginine-rich defensins (RTD-1, BTD-7, Crp-4) and protegrin-1 (PG-1)60 (see Supporting Information, Figure S3). A correlation exists between negative Gaussian curvature and arginine content in structurally similar peptides. HBD-2 and HBD-3, which have ~50% lysines and ~50% arginines, clearly deviate from this curve, although $\langle K \rangle$ is more negative for HBD-3 as compared to HBD-2, consistent with the greater number of arginines in HBD-3.

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Cationic amino acids work differently. We have demonstrated lipid headgroup organization by arginine and lysine generates distinct types of membrane curvature deformations. These
interactions provide an AMP with complementary curvature producing mechanisms. Arginine generates negative Gaussian curvature (positive and negative curvatures along the two perpendicular principal directions), whereas lysine generates negative curvature along one direction only. Mechanisms of curvature generation based on headgroup reorganization are inherently sensitive to headgroup chemistry; therefore, such mechanisms will be conducive to mediating specific peptide—lipid interactions.

On the basis of the above, a series of rules governing the amino acid content of membrane disruptive antimicrobial peptides can be proposed. A decrease in arginine content in a peptide sequence will imply that less negative Gaussian curvature can be generated. This can in principle be compensated for by an increase in both lysine and hydrophobic content. This "exchange rate" concept between arginines and lysines/hydrophobes is illustrated in a comparison between the amino acid content of β-sheet-bonded defensins and that of α-helical AMPs (Figure 4A,B). The average arginine composition of α-helical AMPs is ~50% that of defensins, but the proportion of most hydrophobic amino acids in α-helical AMPs is ~200% that found in defensins (Figure 4A). Likewise, whereas most of the hydrophilic amino acids are found in similar proportions for α-helical AMPs and defensins, the proportion of lysines in α-helical AMPs has drastically increased by ~300% relative to that in defensins (Figure 4B). Thus, there is a unifying conceptual link between α-helical AMPs and β-sheet AMPs: β-sheet AMPs generally rely more on arginines to generate curvature, while α-helical AMPs rely more on lysines and hydrophobes to do so. In both cases, the end result is the generation of saddle-splay curvature.

Within a more general compass, a strong manifestation of this "saddle-splay curvature selection rule" for amino acid content can be observed in the compositions of AMPs. We test this selection rule against 1080 known cationic peptides with activity against Gram positive or Gram negative bacteria.27 Figure 5A shows a plot of N_N/\(N_R\) (the ratio of the number of lysines to the number of arginines) versus % hydrophobic (% of hydrophobic amino acids in peptide, see Experimental Section), in which all AMPs have been binned into seven histograms. A strong increasing trend consistent with the saddle-splay curvature selection rule can be discerned. Figure 5B–D shows a similar analysis of AMP amino acid sequences using more sophisticated measures to quantify peptide hydrophobicity, using three established scales (Kyte–Doolittle,62 Eisenberg consensus,63 Wimley–White biological64), in which all AMPs have been partitioned into 100 bins. A strong positive correlation is observed between hydrophobicity and \(N_N/N_R\). Even though we do not account for effects due to proximate amino acid sequences, peptide conformation in our simple approach, there is strong "exponential-like" dependence between hydrophobicity and \(N_N/N_R\). Even though we do not account for effects due to proximate amino acid sequences, peptide conformation in our simple approach, there is strong "exponential-like" dependence between hydrophobicity and \(N_N/N_R\).

5. CONCLUSIONS

The curvature selection rule is consistent with different mechanisms of barrier disruption such as budding, blebbing, pore formation, and vesiculization. However, many AMPs have additional antimicrobial functions other than permeation. For example, indolicidin,65 buforin,66 and tachyplesin67 are also known to bind internal targets in bacteria such as DNA. Mammalian α- and β- defensins and cathelicidin LL-37 play other important functions in host defense including modulation of the innate immune response and wound repair.10,68 A necessary and sufficient condition for membrane disruption

Figure 5. Demonstration of the exchange between R with K and hydrophobicity based upon 1080 cationic AMPs in the antimicrobial peptide database. (A) By organizing all 1080 AMP sequences into seven histogram bins, a strong positive dependence between the peptide lysine to arginine ratio and peptide hydrophobicity percentage can be observed. More robust calculations of hydrophobicity based upon the widely used Kyte–Doolittle (B), Eisenberg consensus (C), and Wimley–White (D) hydrophobicity scales show the same relationship between the exchange of arginine with lysine and hydrophobicity consistent with the saddle-splay curvature selection rule. See the Experimental Section for details and binning schemes in (B)–(D).
is not a necessary and sufficient condition for full antimicrobial activity, which may involve other mechanisms such as DNA binding. (They may share related amino acid requirements, because both membrane activity and DNA binding require high cationic charge density.) It should be noted, however, that AMPs that kill bacteria by binding to interior elements must first permeabilize the cytoplasmic bacterial membrane to reach the target. Because negative Gaussian curvature is topologically required for pore formation, we expect the saddle-splay curvature selection rule should also apply to nonlytic AMPs provided that membrane permeation is necessary for their function, even if it is not a sufficient condition for full activity.

In a more general context, the saddle-splay curvature selection rule we propose does not strictly restrict the amino acid content of an AMP, because it affects only the arginine, lysine, and hydrophobic residues. Membrane destabilizing AMPs therefore inherently have significant sequence flexibility for beneficial mutations. Consistent with this observation, it is known that AMP amino acid sequences are fairly nonconserved and labile across species. However, when they acquire specific functions, such as Fe homeostasis activity in the hepcidins, their sequence tends to stabilize. That 1080 known AMPs are consistent with the saddle-splay selection rule suggests that they all have some degree of membrane curvature generating activity, but because the selection rule underdetermines the full sequence, there is potential for AMPs to acquire new functions, such as optimized DNA binding.

The saddle-splay curvature selection rule ultimately originates from geometry, so the amino acid composition trends are expected to be general for a broad range of membrane-active peptides. For example, the fusion protein M2 from the influenza virus induces in vitro cell membrane budding and scission. It contains a 17 residue amphiphatic helix with amino acid content comparable to that of AMPs and is capable of inducing budding in GUVs. During virion release, M2 localized to the “neck” of budding virions, the region of maximal negative Gaussian curvature. This suggests other membrane-active peptides and proteins utilize similar strategies to generate curvature in membranes. In summary, we show that a broad range of defensins selectively permeate bacterial membranes by inducing saddle-splay membrane curvature in a manner that depends on target membrane lipid composition. Because the presence of saddle-splay curvature is necessary for membrane-disrupting processes, generation of saddle-splay membrane curvature by AMPs is a versatile mechanism of action. A saddle-splay curvature selection rule that places constraints on the amino acid composition of membrane disruptive peptides is deduced. We believe this rule can guide the rational design of antimicrobials.

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**REFERENCES**


**ASSOCIATED CONTENT**

Supporting Information. SAXS measurements on SUVs used in this study; interaction of β-defensin RTD-1 with pure DOPE lipids and pure DOPG lipids; and SAXS experiments on the cathelicidin protegrin-1 with DOPS/DOPE/DOPC membranes. This material is available free of charge via the Internet at http://pubs.acs.org.

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the full phase diagram of defensin-lipid structures. Pn

the observed re

Pn

a


(32) While coexisting phases (such as lamellar and inverted hexagonal phases) are sometimes observed at specific lipid compositions, the observed reflections cannot be accounted for without the presence of a Pn3m cubic phase. The strong induced formation of cubic phases like the P63m, which are rich in saddle-splay curvature, is a generic feature in the full phase diagram of gemini-lipid structures.


(61) In Figure 4, we exclude the amino acids cysteine, proline, and glycine, which primarily influence AMP tertiary structure and are not directly involved in the cationicity or hydropobicity of AMPs. Also, we consider only arginine and lysine in our exchange rate concept because they constitute the vast majority (89%) of cationic AA found in the 1080 AMPs.