A $^2$H Solid-State NMR Study of Lipid Clustering by Cationic Antimicrobial and Cell-Penetrating Peptides in Model Bacterial Membranes

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ABSTRACT Domain formation in bacteria-mimetic membranes due to cationic peptide binding was recently proposed based on calorimetric data. We now use $^2$H solid-state NMR to critically examine the presence and absence of domains in bacterial membranes containing zwitterionic 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylethanolamine (POPE) and anionic 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylglycerol (POPG) lipids. Chain-perdeuterated POPE and POPG are used in single-component membranes, binary POPE/POPG (3:1) membranes, and membranes containing one of four cationic peptides: two antimicrobial peptides (AMPs) of the $\beta$-hairpin family of protegrin-1 (PG-1), and two cell-penetrating peptides (CPPs), HIV TAT and penetratin. $^2$H quadrupolar couplings were measured to determine the motional amplitudes of POPE and POPG acyl chains as a function of temperature. Homogeneously mixed POPE/POPG membranes should give the same quadrupolar couplings for the two lipids, whereas the presence of membrane domains enriched in one of the two lipids should cause distinct $^2$H quadrupolar couplings that reflect different chain disorder. At physiological temperature (308 K), we observed no or only small coupling differences between POPE and POPG in the presence of any of the cationic peptides. However, around ambient temperature (293 K), at which gel- and liquid-crystalline phases coexist in the peptide-free POPE/POPG membrane, the peptides caused distinct quadrupolar couplings for the two lipids, indicating domain formation. The broad-spectrum antimicrobial peptide PG-1 caused ~40% of the POPE lipids while disordering POPG. The Gram-negative selective PG-1 mutant, IB549, caused even larger differences in the POPE and POPG disorder: ~80% of POPE partitioned into the ordered phase, whereas all of the POPG remained in the disordered phase. In comparison, TAT rigidified POPE and POPG similarly in the binary membrane at ambient temperature, indicating that TAT does not cause dynamic heterogeneity but interacts with the membrane with a different mechanism. Penetratin maintained the POPE order but disordered POPG, suggesting moderate domain separation. These results provide insight into the extent of domain formation in bacterial membranes and the possible peptide structural requirements for this phenomenon.

INTRODUCTION

Lateral phase separation is well recognized for cholesterol- and sphingomyelin-rich lipid membranes that mimic the plasma membrane of eukaryotic cells (1,2). Recently, this domain formation was also proposed for model bacteria membranes containing both zwitterionic and anionic lipids (3). The proposal was based on the observation that bacteria-mimetic 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylethanolamine (POPE)/1-palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylglycerol (POPG) and POPE/cardiolipin membranes containing certain cationic peptides show a phase transition at a high temperature that is similar to the transition temperature of the pure POPE membrane (4,5). In contrast, the phase transition of the peptide-free mixed membranes is broad and occurs at a lower temperature, which is intermediate between the transition temperatures of the two constituent lipids. These data led to the hypothesis that cationic peptides sequester anionic lipids, so that the zwitterionic POPE is segregated into a single-component domain whose physical chemical property is similar to that of the pure POPE membrane. It was further proposed that this charge clustering and domain formation may explain the Gram-selective activity of some cationic antimicrobial peptides (AMPs): Gram-negative bacteria such as Escherichia coli contain significant amounts of both zwitterionic and anionic lipids in their cytoplasmic membranes, whereas the membranes of many Gram-positive bacteria such as Staphylococcus aureus contain predominantly anionic lipids such as POPG and cardiolipin, and therefore cannot form domains (6). This domain formation does not appear to be ubiquitous for all cationic membrane peptides: peptides that cluster anionic lipids appear to have the common features of a high density of positive charges and conformational flexibility, although the number of characterized lipid-clustering peptides is still small.

Indirect evidence of domain formation in model bacterial membranes has also been reported based on small-angle x-ray scattering (SAXS) experiments of POPE-rich membranes containing cationic peptides such as antimicrobial defensins and cell-penetrating peptides (CPPs) such as HIV TAT. When the molar ratios of phosphatidylethanolamine (PE) and anionic lipids are comparable to the bacterial membrane composition, which contains 20–30% anionic lipids, saddle-splay or negative Gaussian curvature was observed (7,8). This Gaussian curvature generation was proposed to be important for membrane processes...
such as pore formation, blebbing, and vesicularization (7). Lower PE content weakens the negative Gaussian curvature, suggesting that the cationic peptides may cause PE enrichment in these model bacterial membranes.

Elucidating the lipid interactions of cationic peptides is important for understanding the structural basis for the actions of AMPs and CPPs (9,10). These Arg-rich peptides share similar amino acid sequences but distinct biological activities. AMPs constitute the primary line of defense by the innate immune system of many multicellular organisms (11,12). They disrupt the barrier function of the lipid membranes of microbial cells by forming membrane pores or by micellizing the lipid membrane. In contrast, CPPs can cross the membrane into cells without damaging the cell membrane, and can do so while carrying macromolecular cargos (13). Although most AMPs are amphipathic molecules with well-defined structures, solid-state NMR (SSNMR) studies showed that two prototypical CPPs, HIV TAT and penetratin, are highly mobile and are both unstructured or low in strand and helix content in the lipid membrane (14,15). This conformational flexibility was proposed to underlie the rapid translocation of CPPs across the lipid membrane, without being stuck in the membrane to cause long-lasting defects (9). NMR distance measurements and two-dimensional correlation spectra showed that Arg-phosphate interactions and Arg-water interactions are common in both AMPs and CPPs (15–17) to stabilize the positive charges in the hydrophobic part of the lipid membrane.

In this work, we use $^2$H SSNMR to investigate whether membrane domains are induced by four cationic AMPs and CPPs, to test the hypothesis that lipid clustering may be one of the mechanisms of action of these peptides. We investigate whether lipid domains that are stable on the millisecond timescale or longer and that are larger than ~30 nm exist in the model bacterial membrane, POPE/POPG (3:1). We measure acyl-chain $^2$H quadrupolar couplings, which reflect the C-H order parameters of POPE and POPG in the binary membrane in the absence and presence of the peptides. If membrane domains are induced by the peptides, the quadrupolar couplings of the lipids in the peptide-poor domain should approach the couplings of the corresponding single-component membrane, and we can quantify the distribution of each lipid in the different domains by simulating $^2$H spectra as a superposition of basis spectra extracted from limiting conditions.

Four cationic peptides are chosen for this study. The $\beta$-hairpin AMP protegrin-1 (PG-1) has been extensively studied using SSNMR (9,10) and molecular dynamics simulations (18,19). This 18-residue hexa-Arg peptide forms oligomeric transmembrane $\beta$-barrels in POPE/POPG (3:1) membranes (20,21). It has nonselective antimicrobial activities against both Gram-negative and Gram-positive bacteria, with minimum effective concentrations in the 1–2 $\mu$g/ml range (22). The second peptide is a PG-1 mutant, IB549, in which all six Arg residues have been replaced by Lys. IB549 is selective against Gram-negative bacteria: it is about twofold more potent than PG-1 against Gram-negative Pseudomonas aeruginosa but 32-fold less active than PG-1 against Gram-positive methicillin-resistant S. aureus (23). Thus, IB549 is a good candidate for investigating whether lipid clustering is correlated with Gram selectivity of AMPs. We also studied two cell-penetrating peptides, HIV TAT and penetratin. The 15-residue TAT peptide contains 6 Args and 2 Lys residues in a contiguous stretch. This highly charged peptide is a random coil in lipid membranes (15) and generates negative Gaussian curvature based on SAXS data (7). For comparison, the CPP penetratin is less mobile and contains $\beta$-turns interspersed with short segments of $\beta$-strands (14). We show that these conformationally and dynamically distinct peptides do not induce large and stable lipid domains at physiological temperature, but at ambient temperature, the two AMPs indeed induce membrane domains enriched in one of the two lipids.

**MATERIALS AND METHODS**

**Membrane sample preparation**

POPE, POPG, 1-palmitoyl-sn-glycerol-3-phosphoethanolamine (d$_{13}$-POPE), and 1-palmitoyl-sn-glycerol-3-[phosphorac-(1-glycerol)] (sodium salt) (d$_{13}$-POPG) were purchased from Avanti Polar Lipids (Alabaster, AL). All peptides were synthesized by Fmoc solid-phase peptide synthesis protocol and purified by HPLC to >95% purity. The amino acid sequences are PG-1 (NH$_2$-RGGRKCTRQKRFQ-KQK-CONH$_2$), TAT(48–60) (GRKKRRQRRRPPQ-CONH$_2$), and penetratin (RQIKIWFQ-NRR-MKWKK-CONH$_2$).

POPE, POPG, and POPE/POPG (3:1) membranes were prepared by dissolving the lipids in chloroform/methanol (1:1) solution, dried under nitrogen gas, redissolved in cyclohexane, and lyophilized overnight. The dry and homogeneous lipid powder was dissolved in 2.5 mL Tris buffer (10 mM, pH 7.5) and freeze-thawed 10 times to obtain uniform vesicles. The peptide was dissolved in 0.5 mL Tris buffer and mixed with the lipid vesicle solutions. Peptide-lipid mixing caused immediate and significant precipitation, indicating strong binding of the peptides to the lipids. The solution was incubated overnight at room temperature, and then centrifuged at 55,000 rpm and 4°C for ~6 h to obtain a homogeneous membrane pellet. The pellet was spun down into a pipette tip, air-dried to a hydration level of 35–40%, and then spun into a 4 mm magic-angle-spinning rotor. All membrane samples were hydrated with regular water containing deuterium spins at their natural abundance of 0.01%. The lack of an isotropic peak in all peptide-free membrane spectra confirms that at the hydration level used, the amount of natural abundance deuterated water was below the detection limit. The peptide/lipid molar ratio was 1:15 for all samples.

**SSNMR experiments**

All SSNMR spectra were measured on a wide-bore Bruker AVANCE 600 MHz (14.1 T) spectrometer operating at 600.13 MHz and 92.12 MHz for $^1$H and $^2$H, respectively. A 4 mm magic-angle-spinning probe was used in the static mode for all $^1$H experiments. $^2$H spectra were measured from 308 to 283 K. Sample temperatures were direct readings from the thermometer positioned near the sample cavity and are estimated to be within 1 K of the true temperature for these static experiments. All samples were stabilized at the desired temperature for at least 1 hour before the spectrum was recorded. $^2$H quadrupolar echo spectra were measured using echo
temperatures (component membrane. The gel-to-LC phase transition mediate between the values of each lipid in its own single-couplings; moreover, these couplings should be inter-meter scale, they should then exhibit similar quadrupolar
If POPE and POPG are homogeneously mixed on the nano-meter scale, they should then exhibit similar quadrupolar
Thus, at a given temperature, the pure POPE membrane has larger couplings than the pure POPG membrane (Fig. 1 a) due to the higher order of POPE lipids. If membrane domains enriched in one of the two lipids are present, the corresponding quadrupolar couplings should then revert to or approach the couplings of the single-component membrane at that temperature. We first show 2H spectra measured at 308 K, near physiological temperature, and then describe the 2H spectra at lower temperatures of 298–283 K.

Fig. 1 shows the 308 K 2H spectra of d31-POPE and d31-POPG in single- and two-component membranes without and with PG-1 and TAT. Four sets of samples are compared: the one-component POPE and POPG membranes, the peptide-bound one-component membranes, the peptide-free POPE/POPG membrane, and the peptide-bound POPE/POPG membrane. Each spectrum is a superposition of the symmetric Pake patterns of all the methylene and methyl groups along the palmitoyl chain. The 2H spectra of the single-component membranes reflect the order parameter distribution along the acyl chain when all lipid molecules undergo the same motions. For simplicity, we use the maximum 2H quadrupolar coupling, read off from the outermost 90° edges of the spectra, to represent the overall

![Graph](image_url)

### RESULTS

#### 2H spectra of d31-POPE and d31-POPG at 308 K

We use 2H quadrupolar couplings of d31-POPE and d31-POPG to probe whether the anionic POPG is clustered by the cationic peptides to form a separate domain from POPE. 2H quadrupolar couplings reflect the time-averaged orientations of the C-H bonds relative to the bilayer normal. Large-amplitude motions of disordered lipids lead to small 2H quadrupolar couplings, i.e., small C-H order parameters. POPE and POPG are homogeneously mixed on the nanometer scale, they should then exhibit similar quadrupolar couplings; moreover, these couplings should be intermediate between the values of each lipid in its own single-component membrane. The gel-to-LC phase transition temperatures (Tm) of undeuterated POPE and POPG are 298 and 271 K, respectively. Deuteration decreases the Tm by a few degrees, as confirmed experimentally below. Thus, at a given temperature, the pure POPE membrane

![Graph](image_url)

**TABLE 1** Maximum 2H quadrupolar couplings (kHz) of lipid membranes in the absence and presence of cationic membrane peptides at 308 K

<table>
<thead>
<tr>
<th>Membrane Type</th>
<th>2H Quadrupolar Coupling (kHz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>d31-POPE</td>
<td>26.2</td>
</tr>
<tr>
<td>d31-POPE + PG-1</td>
<td>26.2</td>
</tr>
<tr>
<td>d31-POPE + TAT</td>
<td>26.2</td>
</tr>
<tr>
<td>d31-POPE/POPG</td>
<td>26.2</td>
</tr>
</tbody>
</table>

![Graph](image_url)
at the end of the acyl chain. In most spectra shown here, the maximum and minimum quadrupolar couplings correlate: larger C2 quadrupolar coupling is associated with larger methyl splitting. No dePaking (26,27) was attempted because the spectra of many peptide-bound membranes do not have sufficient resolution, which may partly result from the presence of intermediate-timescale motion of the acyl chains.

In the absence of any peptide, the pure POPE membrane exhibited a maximum quadrupolar coupling of 36.6 kHz at 308 K. Adding POPG to 25 mol % reduced the POPE couplings significantly, to a maximum value of 30.3 kHz (Fig. 1, a and b), indicating that the low-melting POPG increased the disorder of the high-melting POPE. PG-1 binding to the POPE membrane similarly reduced the maximum coupling to 31.5 kHz (Fig. 1 c), indicating that PG-1 exerted a similar disordering effect as POPG. PG-1 binding to the POPE/POPG membrane reduced the POPE quadrupolar couplings further, to 25.8 kHz (Fig. 1 d). Thus, instead of promoting a more ordered POPE-rich domain with couplings approaching that of the pure POPE membrane, PG-1 increased the POPE disorder, thus ruling out a POPE-rich domain at 308 K.

Compared to POPE, the single-component POPG membrane shows much smaller quadrupolar couplings at 308 K, with a maximum value 23.6 kHz. Mixing with POPE increased the POPG couplings to the same values as the POPE couplings in the binary membrane (Fig. 1 b), indicating unambiguously that the two lipids are homogeneously mixed and exist in a single LC phase at this temperature. PG-1 binding to the POPG membrane moderately decreased the quadrupolar couplings (maximum value 21.1 kHz) compared to the peptide-free POPG membrane. In contrast, PG-1 binding to the POPE/POPG membrane caused a maximum POPG coupling of 24.4 kHz, which is smaller than the coupling of the peptide-free POPE/POPG membrane but higher than the couplings of the POPG membrane with or without PG-1. Instead, the 24.4 kHz coupling is similar to the maximum POPG coupling in the same peptide-bound binary membrane, indicating that PG-1 largely maintained the homogeneous mixing of the two lipids. Therefore, the PG-1-bound POPE/POPG membrane does not show any significant domain formation at 308 K.

TAT shows very different lipid interactions from PG-1 at 308 K. The POPE quadrupolar couplings are similar (~31 kHz) for TAT-bound d31-POPE membrane, peptide-free d31-POPE/POPG membrane, and TAT-bound d31-POPE/POPG membrane (Fig. 1, b, e, and f), indicating that TAT behaves identically to POPG in causing disorder to POPE. In addition, TAT-bound POPG membrane has similar couplings as the pure d31-POPG membrane, and TAT-bound POPE/d31-POPG membrane has similar couplings as the peptide-free POPE/d31-POPG membrane. Taken together, these results indicate that TAT does not perturb either the POPE or POPG dynamics in any of these membranes at 308 K.

These 2H spectra indicate the following lipid mixing situations at 308 K. In the absence of peptides, the POPE/POPG membrane is homogeneously mixed, with similar quadrupolar couplings that differ from the individual lipid’s couplings in its own membrane (see the Supporting Material, Fig. S1 a). Binding of PG-1 and TAT to single-component membranes reduced the POPE couplings significantly and the POPG couplings moderately. Thus, both PG-1 and TAT increased the disorder of the single-component membranes. For the POPE/POPG membrane, PG-1 binding caused similar quadrupolar couplings (24–26 kHz) for the two lipids, both of which are much smaller than the couplings of the peptide-free POPE/POPG membranes. Therefore, PG-1 increased the disorder of both lipids in the binary membrane but did not cause domain separation. TAT also induced similar disorder to POPE and POPG in the binary membrane, but the couplings are unchanged from the peptide-free membranes (Fig. S1 b), thus TAT does not change the membrane disorder at all, in contrast to PG-1.

The 2H spectra of d31-POPG in PG-1-bound membranes and TAT-bound POPG membrane show a pronounced isotropic peak (Fig. 1, c–e). These isotropic peaks are only observed when the rest of the palmitoyl chain shows much weaker couplings, and the intensity envelope excluding the isotropic peak is also higher in the middle of these spectra. These features indicate that the isotropic peak results from the chain-end methyl group, which becomes more disordered after peptide binding, therefore the methyl splitting is no longer resolved but merges into a single narrow peak.

To investigate how general these peptide-lipid interactions are, we also measured the 2H spectra of IB549 and penetratin at 308 K (Fig. 2). IB549 gave rise to similar 2H spectra as PG-1: it caused similar but not identical maximum quadrupolar couplings for POPE (26.9 kHz) and POPG (24.6 kHz) (Fig. 2 b). Penetratin behaves in an intermediate regime between PG-1 and TAT: it caused the same quadrupolar couplings for POPE and POPG in the binary membrane, but both couplings are much smaller than the values of the peptide-free membrane (Fig. 2 d, Table 1). Fig. S1 summarizes the maximum quadrupolar couplings of POPE and POPG of the various lipid membranes at 308 K.

Ambient-temperature 2H spectra of peptide-free lipid membranes

Because lipid motion and membrane disorder depend intimately on temperature, we measured the 2H spectra of the POPE/POPG membrane at lower temperatures of 298 to 283 K. We first describe the 2H spectra of peptide-free membranes at these temperatures. The spectra of the pure POPE membrane (Fig. S2, c–e) are clearly broadened below 293 K, consistent with phase transition of POPE at 298 K.
For the POPE/POPG membrane, the $d_{31}$-POPE spectra at 293 and 287 K are a superposition of LC- and gel-phase spectra (Fig. 3, a and b). We quantified the percentages of POPE in the LC and gel phases by simulating the spectra as a superposition of the 308 K spectrum and the 283 K spectrum of the peptide-free POPE/POPG membrane (Fig. 4). The superposition shows that for the 287 K binary membrane, POPE is distributed in the LC and gel phases at a 16%:84% (±2%) ratio. Increasing the temperature by only a few degrees, to 293 K, inverted the relative LC and gel phase concentrations to 82% for the LC phase and 18% for the gel phase. Thus, POPE partitioning between the LC and gel phases is highly sensitive to temperature in this narrow range. These observations are quantitatively consistent with the known phase diagram of the POPE/POPG membrane (28), which was assembled based on DSC, x-ray scattering, and optical microscopy data. The phase diagram indicates a coexistence temperature range of 291–296 K for a POPG mole fraction of 25%. At 293 K, the phase diagram indicates that the LC- and gel-phase concentrations are 29% and 71% (±2%), respectively, and ~75% of POPE is enriched in the gel phase, whereas POPG distribution is 40%:60% between the LC phase and the gel phase, respectively. The $^2$H-NMR determined POPE distribution in the two phases at 287 K is roughly consistent with the predicted phase composition at 293 K, although the 293 K $^2$H spectrum of POPE is roughly consistent with the 295 K phase composition. The small temperature shift between the $^2$H NMR results and the x-ray and calorimetry-deduced phase diagram likely results from the well-known reduction of the phase transition temperature by lipid chain perdeuteration (29,30).

The $d_{31}$-POPG couplings between 293 and 287 K differ slightly (~3 kHz for the maximum coupling) from the couplings at 308 K (Fig. S2). If we neglect this difference, the 293 K $d_{31}$-POPE/POPG spectrum of the POPE/POPG membrane then corresponds to a superposition of 38% LC phase and 62% gel phase, which is in excellent agreement with the phase-diagram prediction of POPG distribution in the two phases (28).

**Ambient-temperature $^2$H spectra of peptide-bound POPE/POPG membranes**

The effects of PG-1 and IB549 binding on the lipid mobility of the binary membranes at ambient temperatures are shown in Fig. 3 and Fig. 5. In contrast to the 308 K situation, PG-1 and IB549 changed the POPE and POPG couplings in opposite directions, indicating lipid clustering and domain formation. At 293 and 287 K (Fig. 3), PG-1 increased the gel-phase intensities of $d_{31}$-POPE while decreasing the gel-phase contribution of the $d_{31}$-POPG spectra. The gel-phase fraction of POPE at 293 K is 40%,
up from 18% for the peptide-free binary membrane, although at 287 K, the gel-phase fraction of POPE increased from 84% to 96% (Table 2). Thus, PG-1 increased the order of POPE while increasing the disorder of POPG, indicating that POPE and POPG are preferentially separated. IB549 increased the rigidity of POPE even more strongly than PG-1 while maintaining the POPG disorder (Fig. 5): the gel-phase fraction of POPE is 83% at 293 K, almost twice the amount for the PG-1-bound binary membrane.

Compared to the two AMPs, TAT affected the POPE/POPG membrane qualitatively differently (Fig. 6): between 298 and 287 K, both POPE and POPG became more ordered in the presence of TAT. Spectral deconvolution indicates that the gel-phase fraction of each lipid is within 5–15% of each other in this temperature range (Table 2). Thus, TAT uniformly increased the order of both lipids instead of causing dynamically different membrane domains. Interestingly, penetratin, although also classified as a CPP, differs from TAT in its interaction with the lipids. Between 293 and 287 K, penetratin caused little change to the POPE disorder (Fig. S3, Table 2) but significantly increased the POPG disorder (smaller quadrupolar couplings). Thus, penetratin interaction with POPE and POPG is intermediate between the behaviors of the AMPs and TAT.

**DISCUSSION**

Although domain formation has been extensively studied in cholesterol-containing synthetic membranes that mimic the eukaryotic membrane (31,32), domain formation in bacterial membranes is a relatively recent proposal (3). 2H NMR is an excellent probe of the dynamic heterogeneity of lipid membranes and has been used to study the phase diagram of ternary membranes containing cholesterol (33,34). For the model bacterial membrane studied here, if zwitterionic POPE and anionic POPG are separated into sufficiently large lateral domains with different chain mobilities, the quadrupolar couplings of the two lipids should then differ and should approach the values of their single-component membranes (Fig. 1a). If, however, the domains are small or transient, lateral diffusion of the lipids between the domains will then average the quadrupolar couplings. For a maximum 2H quadrupolar coupling of 20–30 kHz, chemical exchange on the timescale of ~50 μs or shorter will give fast-averaged spectra. Assuming a lipid lateral diffusion coefficient $D_L$ of $5 \times 10^{-8} \text{ cm}^2/\text{s}$, this timescale translates to a domain radius ($r$) of ~30 nm based on the diffusion equation $\langle r^2 \rangle = 4D_L\tau$. The domains must be smaller than ~30 nm to observe averaged quadrupolar couplings.
TABLE 2  Mole fractions of d31-POPE and d31-POPG in the disordered and ordered phases of the POPE/POPG membrane from 298 to 287 K, obtained from deconvolution of the 2H NMR spectra

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Membrane</th>
<th>Disordered phase</th>
<th>Ordered Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>298 K</td>
<td>d31-POPE/POPG</td>
<td>~100%</td>
<td>~0%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + TAT</td>
<td>65%</td>
<td>35%</td>
</tr>
<tr>
<td></td>
<td>POPE/d31-POPG</td>
<td>~100%</td>
<td>~0%</td>
</tr>
<tr>
<td></td>
<td>POPE/d31-POPG + TAT</td>
<td>50%</td>
<td>50%</td>
</tr>
<tr>
<td>295 K</td>
<td>d31-POPE/POPG</td>
<td>~100%</td>
<td>~0%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + TAT</td>
<td>23%</td>
<td>77%</td>
</tr>
<tr>
<td></td>
<td>POPE/d31-POPG</td>
<td>~100%</td>
<td>~0%</td>
</tr>
<tr>
<td></td>
<td>POPE/d31-POPG + TAT</td>
<td>21%</td>
<td>79%</td>
</tr>
<tr>
<td>293 K</td>
<td>d31-POPE/POPG</td>
<td>82%</td>
<td>18%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + PG-1</td>
<td>60%</td>
<td>40%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + IB549</td>
<td>17%</td>
<td>83%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + TAT</td>
<td>~0%</td>
<td>~100%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + penetratin</td>
<td>82%</td>
<td>18%</td>
</tr>
<tr>
<td></td>
<td>POPE/d31-POPG</td>
<td>38%</td>
<td>62%</td>
</tr>
<tr>
<td></td>
<td>POPE/d31-POPG + PG-1</td>
<td>~100%</td>
<td>~0%</td>
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<td></td>
<td>POPE/d31-POPG + TAT</td>
<td>7%</td>
<td>93%</td>
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<td></td>
<td>POPE/d31-POPG + penetratin</td>
<td>71%</td>
<td>29%</td>
</tr>
<tr>
<td>287 K</td>
<td>d31-POPE/POPG</td>
<td>16%</td>
<td>84%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + PG-1</td>
<td>4%</td>
<td>96%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + IB549</td>
<td>4%</td>
<td>96%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + TAT</td>
<td>~0%</td>
<td>~100%</td>
</tr>
<tr>
<td></td>
<td>d31-POPE/POPG + penetratin</td>
<td>5%</td>
<td>95%</td>
</tr>
<tr>
<td></td>
<td>POPE/d31-POPG</td>
<td>~0%</td>
<td>~100%</td>
</tr>
<tr>
<td></td>
<td>POPE/d31-POPG + penetratin</td>
<td>8%</td>
<td>92%</td>
</tr>
</tbody>
</table>

couplings between the two domains. In other words, if different quadrupolar couplings are observed for the two domains, the domains must be much larger than ~30 nm. Although this size estimate depends on the value of $D_L$, diffusion coefficients in general differ by less than an order of magnitude among divergent lipid membranes. When the two domains are a gel phase and a fluid phase with different $D_L$ values, which can in principle slow down lipid diffusion across the phase boundary, the domain size still cannot be much smaller than several tens of nanometers, because dynamic differences and diffusion coefficient contrasts cannot be maintained across very small domains. Indeed, experimental evidence suggests that the coexistence of multiple phases in a membrane does not cause significant changes of the diffusion coefficients for domain sizes $< ~200$ nm (35), and the low diffusion coefficients measured for gel-phase lipids came from micron-sized membrane domains (36).

The $^2$H NMR spectra shown here indicate that near the human physiological temperature of 308 K, the two AMPs caused very similar but not identical quadrupolar couplings for POPE and POPG, whereas the two CPPs induced identical couplings for the two lipids in the binary membrane (Figs. 1 and 2). The POPE couplings in the binary membrane are 6–10 kHz smaller than the couplings of the pure POPE membrane and the peptide-bound POPE membrane (Table 1), indicating the absence of a large POPE-enriched domain at 308 K. For PG-1 and IB549, the maximum quadrupolar couplings are 1.4 and 2.3 kHz larger for POPE than POPG. These differences are larger than the 0.4 kHz difference for the peptide-free binary membrane, suggesting that the two AMPs cause a small degree of domain separation. However, the main conclusion at 308 K is that the four cationic peptides induce either no or very minor domain separation in the POPE/POPG membrane.

The fact that the two AMPs increased the disorder of POPE and POPG at 308 K is consistent with the membrane-disruptive effect of these $\beta$-hairpin AMPs. HIV TAT did not affect the membrane disorder, which is consistent with its random coil nature (15). In comparison, penetratin increased the membrane disorder, similar to the two AMPs. This may be attributed to the fact that penetratin, unlike TAT, shows $\beta$-turn and short $\beta$-strand conformations (14,37), and the presence of such H-bonded conformations may facilitate the creation of membrane disorder.

The domain phenomenon changes completely at ambient temperatures of 293–287 K. Both AMPs increased the order of POPE while disordering POPG. Spectral deconvolution indicates that LC and gel phases coexist in this temperature range for the peptide-free POPE/POPG membrane, with the two lipids partitioning into the two phases according to the phase diagram: POPE has only a small gel-phase content (18%) at 293 K but becomes predominantly gel phase at 287 K, whereas POPG is ~60% gel phase at 293 K but 100% gel phase at 287 K (Table 2). PG-1 binding increased the gel-phase content of POPE to 40% at 293 K while keeping POPG predominantly disordered. Thus, PG-1 induced membrane domains at these ambient temperatures: the disordered domain contains a mixture of POPE and POPG, whereas the ordered domain consists exclusively of POPE. Relative to the total lipid content, in the PG-1-bound POPE/POPG membrane, all 25% POPG and 45% POPE reside in the disordered phase, whereas the remaining 30% POPE is separated into the gel phase. A POPE gel phase may not be directly involved in pore formation, because the ordered acyl chains should reduce the negative curvature strain of this lipid compared to the LC phase. However, the existence of significant gel phase may inhibit the function of biological membranes in other ways, for example by impeding protein lateral diffusion.

The Gram-negative-selective AMP, IB549, caused even stronger counter-directional changes to the POPE and POPG chain dynamics. At 293 K the gel-phase content of POPE increased to 83% (Fig. 5, Table 2), which is double that of the PG-1-bound membrane, whereas POPG remained similarly disordered as before. The $^2$H spectra of the POPE/d31-POPG membrane in the presence of both IB549 and PG-1 show modestly smaller couplings (Fig. 5 a, Fig. 5 a) compared to the high-temperature spectra of the peptide-free membranes, thus exact deconvolution is not possible. However, the small quadrupolar couplings qualitatively indicate the high disorder of POPG, consistent
with the notion that POPG preferentially interacts with IB549 and PG-1. Taken together, these data indicate that the Gram-negative antimicrobial activity of IB549 is indeed correlated with its stronger lipid-clustering ability. The structural cause for this stronger lipid clustering may be that the Lys ammonium group, due to its three-site jump motion, is less able to form stable H-bonds with lipid phosphate groups, thus it may predominantly interact with the anionic POPG. In contrast, the Arg variant of AMPs can interact with both POPG and POPE well, thus causing less domain separation.

Compared to the two AMPs, TAT affected the order of the binary membrane differently at ambient temperatures. It increased the gel-phase fraction of both POPE and POPG (Fig. 6), and the gel phase appeared at higher temperatures (298 K) than PG-1- and IB549-bound membranes. Thus, although TAT showed almost no effect to the membrane fluidity at 308 K, it exerted a uniform ordering effect on the binary membrane below 298 K, without causing domains. The molecular mechanism for this uniform ordering is still unclear. A hypothesis proposed by Wong and co-workers (7) based on SAXS data of CPPs is that the conformationally flexible and Arg-rich CPPs can efficiently cross-link multiple lipid headgroups through multidentate hydrogen bonds between the Arg guanidinium and lipid phosphate groups. In principle, this mechanism should not distinguish between POPE and POPG, and hence may explain the similar rigidification of both lipids. Direct distance measurement between Arg and phosphates by SSNMR showed the existence of short guanidinium-phosphate distances in many Arg-rich cationic peptides (9,15,17,37–39), although these data did not directly indicate the number of phosphate groups bound to each guanidinium. The second CPP, penetratin, interacts with the lipids in a more analogous fashion to the two AMPs: it enhanced the dynamic disorder of POPG while exerting relatively weak effects on the POPE order, suggesting that the penetratin-bound membrane may have moderate separation of POPE and POPG.

Taken together, these ²H NMR data indicate that lipid clustering in the model bacterial membrane is a partial phenomenon that occurs within a narrow temperature range. This temperature range is well below the human physiological temperature but is near the temperature at which the POPE/POPG membrane naturally shows coexisting phases. Between 293 and 287 K, the two AMPs clustered and disordered the anionic POPG while ordering the zwitterionic POPE. The cationic peptides most likely partitioned into the disordered phase, which also contains POPE, whose mole fraction depends on the peptide and the temperature. Fig. 7 depicts the membrane domains induced by PG-1 at 293 K, the structure of the peptide and the proposed peptide-lipid interactions in the disordered phase (9,10). If we make the reasonable assumption that the disordered phase also has higher curvature, PG-1 most likely then partitions into the disordered phase because previous lipid-peptide distance measurements indicate that PG-1 is located in the high-curvature toroidal pore defects of the membrane (17).
Fig. 7 shows POPE is not exclusively located in the ordered phase but is also present in the disordered phase, which may help to stabilize the curvature of the toroidal pore due to the intrinsic negative curvature of POPE. This lateral domain separation occurs around 293 K for PG-1 and at a slightly higher temperature (~295 K) for IB549. Above these temperatures, the membrane becomes homogeneous, without coexisting LC and gel phases or separate POPE and POPG domains.

These results suggest that the mammalian AMPs studied here, which must act near the physiological temperature of 308 K, do not use domain formation as a significant mechanism against bacterial membranes. However, for AMPs produced by amphibians such as frogs, whose body temperature is mainly controlled by the ambient temperature, the intrinsic ability of POPE and POPG to phase separate at ~293 K may allow these peptides to cluster POPG and induce a separate POPE domain, thus destroying the barrier function of the membranes of Gram-negative bacteria. Further experiments on other AMPs, especially of nonmammalian origin, will be interesting to test this hypothesis. Compared to AMPs, the conformationally flexible HIV TAT exerts little effect on membrane disorder at 308 K, which is consistent with the membrane translocation function of the peptide, but TAT uniformly orders POPE and POPG at lower temperatures, which may reflect the fact that the randomly positioned Arg residues in this unstructured peptide may interact equally well with the phosphate groups of both neutral and anionic lipids.

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