

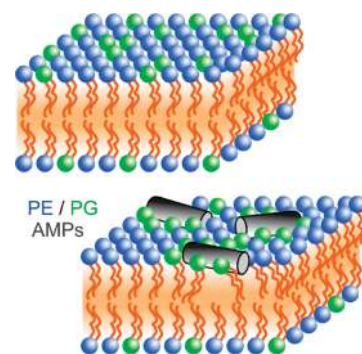
Antimicrobial Peptides Induce Growth of Phosphatidylglycerol Domains in a Model Bacterial Membrane

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ABSTRACT We performed microsecond long coarse-grained molecular dynamics simulations to elucidate the lateral structure and domain dynamics of a phosphatidylethanolamine (PE)/phosphatidylglycerol (PG) mixed bilayer (7/3), mimicking the inner membrane of gram-negative bacteria. Specifically, we address the effect of surface bound antimicrobial peptides (AMPs) on the lateral organization of the membrane. We find that, in the absence of the peptides, the minor PG fraction only forms small clusters, but that these clusters grow in size upon binding of the cationic AMPs. The presence of AMPs systematically affects the dynamics and induces long-range order in the structure of PG domains, stabilizing the separation between the two lipid fractions. Our results help in understanding the initial stages of destabilization of cytoplasmic bacterial membranes below the critical peptide concentration necessary for disruption, and provide a possible explanation for the multimodal character of AMP activity.

SECTION Surfactants, Membranes



Linear α -helical antimicrobial peptides (AMPs), components of the innate immune systems of animals and plants, have prompted growing attention over the past decades as feasible candidates for the treatment of resistant pathogenic bacteria.¹ In order to work against Gram-negative bacteria, AMPs have to pass through the outer lipopolysaccharide membrane, the peptide-glycan layer, and disturb the stability of the inner lipid membrane.¹ Understanding the latter process at the molecular level is beneficial for designing efficient and selective agents that can be used for medical purposes instead of conventional antibiotics. The standard model system in experimental and computational studies of the inner bacterial membrane is a mixed phosphatidylethanolamine (PE)/phosphatidylglycerol (PG) bilayer^{2,3} with an approximate 7/3 ratio, mimicking the inner membrane of *Escherichia coli*. Since this system contains a major (PE) and minor (PG) component, it is reasonable to assume microphase separation in the membrane plane, resulting in the formation of PG domains surrounded by the PE medium. Recently, the formation of stable lipid domains in monolayers of lipids from *E. coli* extracts has been shown using atomic force microscopy.⁴ In addition, anionic bacterial lipids with high intrinsic curvature (such as cardiolipin) are known to lead to the formation of finite-sized lipid domains in a curvature-dependent manner.⁵ Clustering of anionic lipids (PG and cardiolipin) in bacterial membranes has been proposed to play an important role in the functioning

of membrane proteins, cell division, and the action of antimicrobial agents.^{6,7} Direct experimental evidence for peptide-induced domain formation in bacterial-membrane-mimetic supported bilayers was recently reported.⁸

We performed coarse-grained (CG) molecular dynamics (MD) simulations of mixed dioleoyl-PE/dioleoyl-PG (70/30%) bilayers in order to elucidate with molecular resolution the structure of the minor anionic phase in the bacterial membrane. We then addressed the effect of AMP binding on lipid clustering. For this purpose, we chose a recently isolated and well-characterized linear α -helical AMP from the Laticin family, Ltc1.^{9,10} Ltc1 is cationic with an overall charge of +10. The peptide/lipid ratio in our simulations is 1:20. This ratio is slightly lower than the critical concentration of the peptide that causes prominent destabilization of PE/PG mixed bilayers.¹⁰ To model the peptides and lipids, we used the CG Martini force field,^{11,12} which represents on average four atoms and associated hydrogens by an effective interaction site. The model has been applied previously in a number of membrane/peptide studies.¹³ According to NMR data, Ltc1 forms an extended α -helix in membrane-mimicking media.¹⁰

Received Date: August 16, 2010

Accepted Date: October 5, 2010

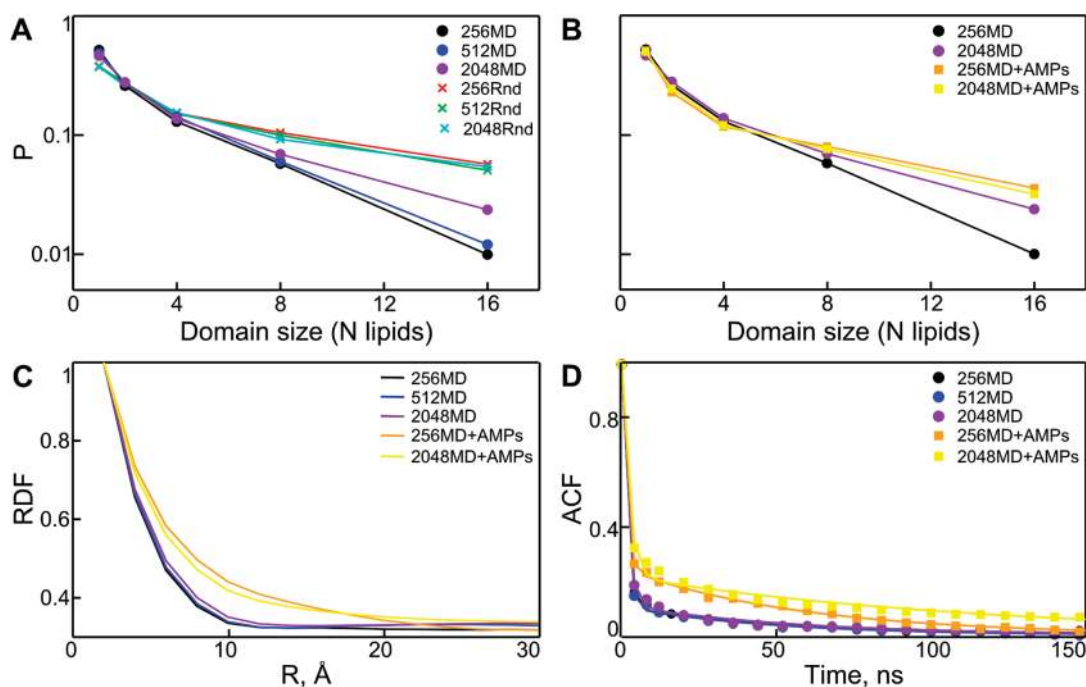


Figure 1. The effect of binding of AMPs on PG domain formation in a PE/PG bilayer. (A,B) Log-normal plots of PG domain size distributions. (C) 2D radial distribution functions (RDFs) of “bilayer maps”. (D) Time autocorrelation functions (ACFs) of PG density maps (with double exponential fits). Systems of 256, 512, and 2048 lipids, either from MD simulations (256MD, 512MD, 2048MD) or randomized bilayers (256Rnd, 512Rnd, 2048Rnd), and systems with 256 or 2048 lipids and 12 or 96 AMPs (256MD+AMPs, 2048MD+AMPs), respectively, are compared.

In our MD simulations, we thus constrained the conformation of the AMPs to be helical. We studied five systems: three PE/PG mixtures composed of 256, 512, and 2048 lipids, respectively (labeled 256MD, 512MD, 2048MD), and systems of either 256 or 2048 lipids with 12 or 96 Ltc1 AMPs, respectively (256MD+AMPs, 2048MD+AMPs). All systems contain ~ 40 waters/lipid and sodium ions to neutralize the PG lipids. For each of the systems, a $4 \mu\text{s}$ simulation (effective time¹⁴) was performed after an initial equilibration phase. For the lipid/peptide systems, the peptides were observed to embed themselves into the bilayer interface equally for both leaflets, staying almost parallel to the surface without creating structural defects such as pores. To analyze the clustering propensity of PG lipids, the following procedure was used. First, the solvent-accessible surface (SAS) of each bilayer snapshot was computed using the PLATINUM software.¹⁵ Surface points were then classified according to lipid type (PE or PG). The SAS surface points were subsequently interpolated to a two-dimensional (2D) grid at 2 \AA resolution (“bilayer maps”), followed by a neighbor-search procedure¹⁶ for domain delineation. Each bilayer leaflet has been treated separately. All quantities reported in this paper are averages over both leaflets. More details about the simulation setup, equilibration, and analysis are given in the Supporting Information (SI).

The results of the simulations with respect to the structural and dynamical behavior of the anionic lipids are shown in Figure 1. In Figure 1A the PG domain size distribution of the mixed bilayer is compared to that expected for a randomized mixture. The randomly assembled bilayer (with the same geometrical parameters and lipid composition as the MD one)

provides the baseline domain size distribution with which the other cases are compared. The normalized frequencies of PG domain sizes decline exponentially with increasing domain size. This is in agreement with the well-known scaling behavior of domain sizes below the percolation threshold.¹⁷ However, we observe a difference between randomized and MD bilayers, with the domain size distribution of the randomized system biased toward larger domains. This effect is present for all systems tested, and implies that PG lipids in MD bilayers try to disperse themselves into smaller clusters. We attribute this effect to the electrostatic repulsion between the negatively charged PG lipids. These unfavorable interactions are partially compensated by preferential binding of sodium ions to PG head groups (Figure S1, SI).

In Figure 1B we compare the PG domain size distribution in the bilayer in the absence or presence of AMPs. Binding of AMPs to the bilayer surface results in a more heavy-tailed domain size distribution, suggesting aggregation of PG lipids. Although the effect seems to decrease for the larger system, the increased clustering propensity of PG lipids induced by AMPs is confirmed by the 2D radial distribution function (RDF, Figure 1C). The faster decline of the RDF of AMP-free systems, in comparison to AMP-containing bilayers, reveals that the binding of AMPs to the bilayer surface induces long-range order in the PG domains and increases their average size. In addition, the AMPs are found to alter the dynamic behavior of the domains. We quantify this by spatially averaged time autocorrelation functions (ACFs) of the 2D bilayer maps (Figure 1D). The difference between AMP-free and AMP-containing systems seems to be robust over system sizes. The presence of

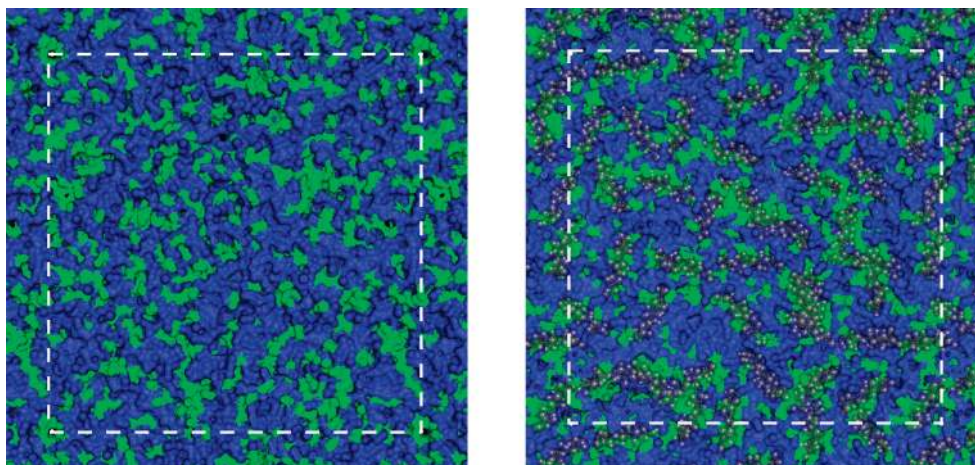


Figure 2. Visualization of AMP-induced PG domains. Snapshots of the surface of a 2048-lipid PE/PG bilayer without (left) and with (right) Ltc1 peptides are shown (the top view). The bilayer surface is colored according to lipid type (PE blue, PG green). Peptides are depicted with semitransparent gray spheres. The degree of transparency corresponds to the distance from the bilayer surface. Ions and water molecules are not shown for clarity. Only one of the bilayer leaflets is rendered. The simulation box is shown with a dashed rectangle.

AMPs systematically leads to longer correlation times. The ACFs can be fitted with double exponentials, yielding two characteristic relaxation times, τ_1 and τ_2 (see SI for details). All three peptide-free systems have comparable relaxation times with $\tau_1 = 1.6 \pm 0.1$ ns and $\tau_2 = 59 \pm 7$ ns on average. The relative contributions of the fast/slow components, given by the ratio of exponential prefactors, $R = A/B = 0.90 \pm 0.01/0.1 \pm 0.01$. We observe that in the small system AMPs do not strongly affect the relaxation times ($\tau_1 = 1.3$ ns and $\tau_2 = 57$ ns in the 256MD+AMPs system). The relative contribution of the slow component, however, increases 2-fold in the presence of AMPs ($R = 0.75/0.25$). The effect of peptides in the large system is even more pronounced. Apart from a similar redistribution between slow and fast components ($R = 0.78/0.22$), the AMPs also cause a 2-fold increment of τ_2 as compared to the peptide-free systems ($\tau_1 = 2.2$ ns, $\tau_2 = 113$ ns in the 2048MD+AMPs system). If we assume that in the peptide-free system the sodium counterions stabilize the scattered PG domains, we can attribute the fast component τ_1 to the self-diffusion of single, dispersed PG lipids and the slow component τ_2 to the movement of entire domains. The increased contribution of the slow component and of the relaxation time τ_2 in the systems with AMPs thus reflects a tendency to form more stable and larger PG domains (cf. Figure 1B, C). It has previously been observed in fully atomistic MD simulations that the “trapping” of lipids by a membrane-active peptide results in their effective freezing.¹⁸ Binding of counterions has a similar effect.¹⁹

Visualization of the domains, presented in Figure 2, emphasizes the existence of PG clusters correlated with the presence of AMPs (Figure 2, right). The situation in the absence of peptides is different (Figure 2, left); here, the PG lipids show a preference to be dispersed into small clusters, presumably due to electrostatic repulsion. Taken together, the results of our simulations provide evidence for a peptide-induced clustering and long-range ordering of PG lipids in mixed PG/PE bilayers mimicking bacterial cell membranes. Interestingly, the peptide-induced clustering is coupled to a

lipid-induced peptide clustering. Analysis of the last 100 ns of 256MD+AMPs and 2048MD+AMPs trajectories gives the following average number and standard deviation of peptides coexisting in a single PG-cluster: 1.6 ± 1.0 and 2.8 ± 2.8 , respectively. Although there is still a finite-size effect, the tendency is clear: interactions of cationic AMPs with anionic PG lipids facilitate peptide oligomerization, likely caused by charge neutralization.

Our results help understand the possible initial stages of destabilization of the cytoplasmic bacterial membrane. On one hand, the preferential interaction between PGs and AMPs leads to a local increase in the peptide surface density, a favorable condition for the formation of pores.²⁰ On the other hand, the recruitment of PGs into microdomains will affect the structural and dynamic properties of the bacterial membrane on a global level. For instance, the bacterial membrane contains a large fraction of PE lipids, which are not able to form stable bilayers on their own. Thus, “purification” of PE as a result of PG “extraction” induced by AMPs would lead to reduced membrane stability. In addition, growing phase boundary defects might be responsible for the increase of bilayer permeability.⁶ Finally, the peptide-induced reorganization of lipids in the membrane could also affect the function of other membrane components, such as membrane proteins. This may lead to a disorganization of metabolic processes in the cell, hampering cell growth and/or proliferation. Such disturbance of the bacterial metabolism, rather than their immediate killing (“sand in a gear box” mechanism) was recently reported for synthetic α -helical peptides.²¹ Peptide-induced changes in the physical-chemical properties of anionic microdomains in the bacterial membrane could be one possible molecular mechanism to explain this multimodal character of AMP activity in bacteria.

SUPPORTING INFORMATION AVAILABLE A table with details of the MD protocols, details of the analysis procedure, and a figure showing the RDF for lipid polar heads and sodium counterions. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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ACKNOWLEDGMENT This work was supported by EMBO (Grant ASTF 85.00-2008), by the Russian Ministry of Science and Education (MK-69.2008.4, MK-8439.2010.4), by the Russian Foundation for Basic Research, and RAS Programs. R.R. was financed by a grant from the Swiss SystemsX.ch initiative, evaluated by the Swiss National Science Foundation (Grant LipidX-2008/011) to I.F.S.

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