Magainin 2 and PGLa in Bacterial Membrane Mimics I: Peptide-Peptide and Lipid-Peptide Interactions

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ABSTRACT We addressed the onset of synergistic activity of the two well-studied antimicrobial peptides magainin 2 (MG2a) and PGLa using lipid-only mimics of Gram-negative cytoplasmic membranes. Specifically, we coupled a joint analysis of small-angle x-ray and neutron scattering experiments on fully hydrated lipid vesicles in the presence of MG2a and L18W-PGLa to all-atom and coarse-grained molecular dynamics simulations. In agreement with previous studies, both peptides, as well as their equimolar mixture, were found to remain upon adsorption in a surface-aligned topology and to induce significant membrane perturbation, as evidenced by membrane thinning and hydrocarbon order parameter changes in the vicinity of the inserted peptide. These effects were particularly pronounced for the so-called synergistic mixture of 1:1 (mol/mol) L18W-PGLa/MG2a and cannot be accounted for by a linear combination of the membrane perturbations of two peptides individually. Our data are consistent with the formation of parallel heterodimers at concentrations below a synergistic increase of dye leakage from vesicles. Our simulations further show that the heterodimers interact via salt bridges and hydrophobic forces, which apparently makes them more stable than putatively formed antiparallel L18W-PGLa and MG2a homodimers. Moreover, dimerization of L18W-PGLa and MG2a leads to a relocation of the peptides within the lipid headgroup region as compared to the individual peptides. The early onset of dimerization of L18W-PGLa and MG2a at low peptide concentrations consequently appears to be key to their synergistic dye-releasing activity from lipid vesicles at high concentrations.

SIGNIFICANCE We demonstrate that specific interactions of the antimicrobial peptides MG2a and PGLa with each other in POPE/POPG bilayers lead to the formation of surface-aligned parallel dimers, which already provide, at low peptide concentrations, the nucleus for the peptides' well-known synergistic activity.

INTRODUCTION

The steady increase of antibiotic resistance of pathogenic bacteria, combined with the decline of approved antimicrobial agents, is considered to be a severe threat to global health. In view of these developments, considerable research efforts have been devoted to understanding the mode of action of antimicrobial peptides (AMPs), considered as an alternative for the development of novel antibiotics. AMPs are effector molecules of the innate immune system whose main targets are bacterial membranes. AMPs kill bacteria within minutes, which makes it more difficult for bacteria to develop resistance mechanisms (for review, see, e.g., (1,2)). Applying diverse biophysical techniques on lipid-only membrane mimics, several interaction models have been conceived for AMPs (1,3,4). Pore formation is arguably the most widely discussed membrane disruptive mechanism.

Choosing a specific membrane mimic requires a delicate balance between experimental or computational tractability and physiological relevance. This leads to a search for the minimum realistic lipid mixture that yields a similar response to AMPs as in live bacteria. Phosphatidylethanolamine, phosphatidylglycerol, and cardiolipin represent the main lipid components of Gram-negative cytoplasmic membranes. Yet, we demonstrated previously that lipid bilayers composed of palmitoyl-oleoyl-phosphatidylethanolamine

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POPE) and palmitoyl-oleoyl-phosphatidylylglycerol (POPG) (molar ratio: 3:1) respond similarly to PGLa and magainin 2-amide (MG2a), in agreement with their antimicrobial activity toward Escherichia coli K12 (5), despite the lack of cardiolipin or other cell wall components. Hence, POPE/POPG (3:1 mol/mol) bilayers appear to be valid first-order mimics for biophysical studies on the activity of AMPs with Gram-negative bacteria.

Studying the activity of PGLa and MG2a is of specific interest because of their synergistic effects, first described by Matsuzaki and co-workers (6–8). Equimolar mixtures of the two peptides reduced the minimum inhibitory concentration of the individual peptides in E. coli K12 by about one order of magnitude. The molecular mechanism of the observed synergy remains controversial, however. Originally, synergism was associated with the formation of a transmembrane pore with a 1:1 peptide stoichiometry (6–8). In contrast, solid-state NMR experiments from the Ulrich and Bechinger groups demonstrated that MG2a never adopts a transmembrane topology and that PGLa may insert perpendicularly in the presence of MG2a into the membrane only in phosphatidylcholine-enriched or short-chain disaturated phosphatidylethanolamine-enriched bilayers, respectively (9–13). None of these latter lipids are of significance for Gram-negative cytoplasmic membranes, however.

The dependence of MG2a-mediated PGLa insertion into bilayers was attributed to the intrinsic lipid curvature, which leads to a tight packing of the bilayer’s polar-apolar interface in the case of cone-shaped lipids such as POPE and leads to a perturbation of membrane structure, which is more pronounced than in the case of noninteracting individual peptides. The resulting remodeling of membrane structure thus appears as a precursor to synergistic dye release at higher peptide concentrations.

**MATERIALS AND METHODS**

**Lipids, peptides, and chemicals**

POPE and POPG were purchased from Avanti Polar Lipids (Alabaster, AL; purity >99%) as powder and used without further purification. L18W-PGLa (GMASKAGAIAGKIAKV AWKAL-NH2) and MG2a (GIGKFLH SAKKFGRAGVEMNS-NH2) were obtained in lyophilized form (purity >95%) from PolyPeptide Laboratories (San Diego, CA). Deuterium oxide (purity 99.8 atom %) and HEPES (purity >99.5) were purchased from Carl Roth (Karlsruhe, Baden-Württemberg, Germany). All other chemicals were obtained from Sigma-Aldrich (Vienna, Austria) in pro analysis quality.

**Sample preparation**

Lipid thin films were prepared by mixing appropriate amounts of lipid stock solutions to obtain samples composed of POPE:POPG (3:1, mol/mol), followed by solvent evaporation under a nitrogen stream at 35 °C and overnight storage in a vacuum chamber. Dry lipid films were hydrated in 10 mM HEPES containing 140 mM NaCl (pH 7.4). For neutron experiments, the H2O/D2O ratio in the buffer was varied as detailed below. Hydrated samples were equilibrated for 1 h at 55 °C, followed by eight freeze-and-thaw cycles using liquid N2 and intermittent vortex mixing. Large unilamellar vesicles (LUVs) were obtained by 31 extrusions with a handheld mini extruder (Avanti Polar Lipids) using a 100 nm pore diameter polycarbonate filter. Vessel size and polydispersity were determined via dynamic light scattering using a Zetasizer NANO ZSP (Malvern Instruments, Malvern, United Kingdom). LUVs were again phosphate assayed and mixed with appropriate amounts of peptide stock solution to obtain peptide:lipid (P:L) molar ratios in the range of P:L = 1:400–1:50. LUVs were equilibrated at a given peptide concentration for 4 days before measurement.

**SAXS**

SAXS data were collected at the SWING beamline (Soleil, Saint-Aubin, France) using x-ray photons of wavelength λ = 1 Å and an Eiger 4M detector (Dectris, Baden-Daettwil, Switzerland). Samples were manually
SANS

SANS experiments were performed at KWS-1 (FRM II, Munich-Garching, Germany (16)). Using a two-dimensional scintillation detector, a wavelength of 5 Å (Δλ/λ = 0.1), and sample-to-detector distance (SDD) = 1.21 and 7.71 m allowed us to cover a q-range from 0.005 to 0.42 Å⁻¹. Samples were kept in 1 mm pathlength quartz cuvettes QX-404 (Helmma, Jena, Germany) and equilibrated at 35°C using a circulating water bath. The contrast conditions used were 100, 75, and 50% v/v D₂O/H₂O. Data correction (integration, normalization, and background subtraction) was performed using the software Foxtrot (Xenocs, Sassen-Flagey, France).

Joint SAXS/SANS analysis

For the P:L ratios used in this study, no peptide-mediated aggregation of LUVs was observed, i.e., scattering data did not exhibit any Bragg peak scattering. This allowed us to treat SAXS/SANS data in terms of dilute particle scattering, that is, we were able to neglect any contribution from LUV-LUV or bilayer-bilayer positional correlations. Moreover, we focused our analysis on peptide-induced changes of the membrane structure by analyzing scattering data for q > 0.05 Å⁻¹. In this q-range, overall vesicle size and morphology do not contribute (17), which allowed us to model the scattered intensities as \( I(q) \propto \left| F_B(q) \right|^2 q^2 \), where \( F_B(q) \) is the flat bilayer form factor.

The form factor was derived within the framework of the scattering density profile (SDP) model (18). In brief, the SDP analysis is based on a composition-dependent parsing of membrane structure into quasimolecular fragments whose distribution along the bilayer normal is described in terms of Gaussian-type volume probability functions. Parsing is guided by MD simulations (for a recent review, see, e.g., (19)). The volume probability functions can be easily scaled with the neutron or x-ray scattering lengths of each group, entailing a joint SAXS/SANS analysis that combines the differently contrasted samples into one underlying membrane structure. Analogous strategies have been reported previously; see, e.g., (20–23).

Parsing

Considering previously reported SDP data for POPE (24) and POPG (25) and by using all-atom MD simulations (see below), the lipid part of the membrane structure was parsed into the ethanolamine (ENX), glycerol (PG), phosphate (PO₄), carbonyl glycerol (CG), CH, CH₂, and CH₃ groups (see Fig. 1; Fig. S1). After our previous SDP analysis of coexisting lipid do-

FIGURE 1 Schematic of the applied SDP model. (A) shows the distribution functions of the different quasimolecular groups (see also Fig. S1). The spheres in the overlaid MD simulations represent phosphor atoms. (B) gives a schematic of the unit cell of cross-sectional area \( A_U \), with different contributions from lipid, peptide, and water. To see this figure in color, go online.

where \( V_{LP} \) is the total volume of the lipid-peptide unit cell (Eq. S9); the hydrocarbon chain length \( D_C = z_{CH_2} \); and the distance from the CG to PO₄ groups \( D_{HD} = z_{PO_4} - z_{CG} \).

Data fitting

To obtain robust fitting parameters and estimates for their uncertainties, every data set was fitted 400 times using a genetic algorithm (31) with random starting parameters. This gave us Gaussian-like distributions for each adjustable parameter. We report the average (center of mass) of these distributions; uncertainties were calculated from second moments. For details regarding constraints and cost function, see the Supporting Materials and Methods.

MD simulations

MD simulations were performed using GROMACS version 2016.2 (32,33).
All-atom simulations

Simulation settings

Protein and solvent molecules were described by Amberff99SB-ILDN (34,35) and lipids by Slipids (36,37) force fields. The simulation time step was set to 2 fs. A temperature of 308.15 K was maintained by using a Nosé-Hoover thermostat (38–40), with a coupling constant of 0.5 ps. Two separate coupling groups for protein-lipid and solvent atoms were used. Three-dimensional periodic boundary conditions were applied, and a Parrinello-Rahman barostat (41,42) with semi-isotropic coupling scheme was employed for keeping the pressure at 1 bar with a coupling constant of 2 ps. Long-ranged electrostatic interactions were treated with the particle mesh Ewald method (43) with the real-space cutoff set to 1.2 nm. Lennard-Jones interactions were cut off at 1.2 nm. All bonds were constrained using the LINCS algorithm; long-range dispersion corrections (44) were applied for energy and pressure.

System preparation

Lipid bilayers composed of 192 POPE and 64 POPG molecules were assembled in the xy-plane by distributing lipids equally in both leaflets using the CHARMM-GUI interface (45). The system was hydrated by more than 40 water molecules per lipid, and NaCl ions were added at 130 mM concentration. The initial box dimensions were 8.9 × 8.9 × 8.5 nm. MG2a and L18W-PGLa were prepared in α-helical conformation.

The following starting configurations were considered. System 1: a single peptide was placed into each membrane leaflet in a surface-aligned topology. System 2: a parallel L18W-PGLa/MG2a heterodimer was placed into each membrane leaflet in a surface-aligned topology.

All systems were equilibrated using similar protocols. Firstly, energy minimization was performed using the steepest-descent algorithm. Then, an equilibration with positional restraints on peptide backbone was performed for 60 ns, followed by an equilibration with dihedral restraints to maintain the peptide’s secondary structure. The length of simulations with dihedral restraints were 105 ns (system 1) or 180 ns (system 2). Finally, unrestrained production dynamics simulations were performed for 500 ns.

Coarse-grained simulations

Simulation settings

Computationally efficient coarse-grained simulations were performed using the MARTINI 2.2 force field (46–48) with a simulation time step of 20 fs. A velocity-rescaling thermostat (modified with a stochastic term) (49) was employed with a coupling constant of 1.0 ps to maintain the temperature at 310 K. Protein-lipid and solvent beads were coupled to separate baths to ensure correct temperature distribution. The pressure was kept at 1 bar using a Parrinello-Rahman barostat with semi-isotropic coupling scheme to maintain the system’s behaviour as a function of time. Finally, we performed a biased simulation in which MG2a and L18W-PGLa were restrained in heterodimers via a flat-bottom potential. The potential was applied to the distances between peptide centers of mass for separations larger than 1 nm with a force constant of 1000 kJ mol⁻¹ nm⁻².

System preparation

The membrane was assembled in the xy-plane using the CHARMM-GUI web server (30). The lipid bilayer was composed of 378 POPE and 126 POPG equally distributed lipids in both leaflets. Roughly 30 water beads per lipid were added (a single bead corresponds to four water molecules), and NaCl ions were also added at a concentration of 130 mM. Two P:L ratios—1:42 and 1:21—were considered for MG2a, L18W-PGLa, and MG2a/ L18W-PGLa (1:1 mol/mol), placing an equal number of randomly distrib-uted peptides into each membrane leaflet. Dimerization was derived from analyzing distances between the peptide centers of mass and the peptide termini. If at least two of these distances were smaller than 1 nm, then the peptides were considered to be in a dimer. As a control, we performed an unbiased simulation starting with peptide heterodimers, taking all parallel and antiparallel variants of mutual peptide alignment into account and analyzed the system’s behaviour as a function of time. Finally, we performed a biased simulation in which MG2a and L18W-PGLa were restrained in heterodimers via a flat-bottom potential. The potential was applied to the distances between peptide centers of mass for separations larger than 1 nm with a force constant of 1000 kJ mol⁻¹ nm⁻².

RESULTS

Scattering experiments

Membrane structure and peptide location

We determined the structural response of POPE:POPG (3:1 mol/mol) LUVs to both peptides, including their equimolar mixture (P:L = 1:200, T = 35°C) using a joint analysis of scattering data with a total of four contrasts (SAXS, SANS: 50, 75, 100% D₂O). No vesicle aggregation or formation of multilamellar aggregates occurred at this peptide concentration, as evidenced by the pure diffuse nature of all scattering patterns (Fig. 2; Figs. S2–S4). This enabled us to derive the peptides’ effect on membrane structure in detail using the analysis described in the previous section.

Pure POPE/POPG bilayers serve as a reference system for our study. Our SDP analysis yields 60.56 ± 0.10 Å² for the lateral area per lipid (Table 1); see Fig. S2 for corresponding fits and Table S1 for all parameter values. This value compares well to A_U = 59.8 Å² obtained by molecular-averaging the individual areas per lipid reported for POPE and POPG (24,25), which supports our analysis. Moreover, our model shows a good agreement with the scattered intensities in the presence of peptides (Fig. 2; Figs. S3 and S4), which further supports our assumption of evenly distributed peptides. An equal distribution of peptides in both leaflets is expected to need extended sample equilibration times, such as realized for the data presented here, during which peptides may spontaneously translocate through the membranes.

We note, however, that we cannot fully exclude the presence of asymmetric transleaflet distributions of the peptides. Disentangling leaflet compositional differences is beyond the resolution of the experiments presented here.

All peptides caused significant modulations of membrane structure (Table 1), with L18W-PGLa having the least effect, e.g., the decrease of thickness of the hydrophobic core 2ΔD_C ≈ −0.4 Å and A_U ≈ 0.7 Å². Interestingly, D_B increased slightly. This can be mainly attributed to the increased volume of the unit cell due to contributions from L18W-PGLa (see Eq. 1). For MG2a in turn, no significant changes were observed for D_B, whereas 2ΔD_C and A_U changed about twice as much compared to L18W-PGLa. In the case of the equimolar peptide mixture, A_U ≈ 2.7 Å² is most significant, leading even to a decrease of D_B. Because 2ΔD_C does
not contain contributions from peptide volumes, it consequently is the appropriate parameter to measure membrane thinning. For the studied peptides, membrane thinning follows the order L18W-PGLa/MG2a > MG2a > L18W-PGLa. We note that the thinning observed for L18W-PGLa/MG2a cannot be explained by a simple linear combination of the thinning effects of the individual peptides. In particular, the experimental form factors of POPE/POPG in the presence of L18W-PGLa and MG2a cannot be averaged to yield the form factor observed upon addition of the equimolar mixture, which should be possible if the two peptides were not interacting with each other (Fig. S5). This supports the formation of heterodimers at significantly lower P:L ratios than previously reported (7,9).

It is particularly interesting to relate membrane thinning to the location of the peptides within the bilayer (Table 1). L18W-PGLa is positioned slightly below and MG2a slightly above the glycerol backbone (CG group), respectively, as observed by comparing \( z_p \) and \( D_C \). The peptide mixture behaved similar to MG2a, with a preferential location just above the lipid backbone. Thus, pronounced thinning effects are observed for peptides located further away from the membrane center, as well as for L18W-PGLa/MG2a, presumably because of dimer formation.

Finally, we present results for the distance between CG and the PO4 groups, \( D_{\text{HD}} \). Because of the rotational flexibility of the lipid headgroup, this value is a measure of the average headgroup tilt projected on to the normal of the lipid bilayer. That is, the lowest \( D_{\text{HD}} \) found in the presence of MG2a indicates that the lipid headgroups are more tilted toward the membrane than for L18W-PGLa/MG2a with the largest \( D_{\text{HD}} \)-value.

**Effect of temperature and peptide concentration**

We varied the temperature in the range from 35 to 50°C to see whether the two peptides and their mixture induce specific changes to membrane structure (Figs. S6–S8). Additionally, we increased the peptide concentration, ensuring that Bragg peaks resulting from vesicle aggregate changes do not dominate the scattered intensities. In the case of MG2a and L18W-PGLa, this allowed us to go up to P:L = 1:50 (Figs. S7 and S8). For the 1:1 peptide mixture, we observed the onset of multilamellar vesicle (MLV) formation (indicated by a low intensity peak) already at P:L = 1:200 (Fig. S9). We thus restricted our analysis to P:L = 1:400–1:200 for the equimolar mixture. Peptide-induced formation of coupled bilayers will be discussed in the subsequent study.

Compared to the joint SAXS/SANS data discussed above, the performed temperature-dependent SAXS experiments provided limited structural resolution. Nevertheless, relative structural changes of structural parameters such as \( A_U \) as a function of temperature can be retrieved reliably (Figs. S10 and S11) using reported lipid-volume temperature dependencies (24,25). Fig. 3 shows the thermal expansion of the area per unit cell for POPE/POPG bilayers (\( \Delta A_U/\Delta T \))

**TABLE 1 Effect of Magainins on the Structure of POPE/POPG (3:1 mol/mol) Bilayers**

| Sample            | \( A_U \) [Å²] | \( D_A \) [Å] | \( D_C \) [Å] | \( z_p \) [Å] | \( D_{\text{HD}} \) [Å] |
|-------------------|----------------|--------------|--------------|--------------|------------------|
| POPE/POPG         | 60.56 ± 0.10   | 39.15 ± 0.06 | 15.34 ± 0.02 (15.10) | –            | 4.19 ± 0.05      |
| + L18W-PGLa       | 61.26 ± 0.13   | 39.49 ± 0.08 | 15.16 ± 0.03 (15.06) | 15.75 ± 0.44 | 4.07 ± 0.06      |
| + MG2a            | 62.00 ± 0.09   | 39.17 ± 0.06 | 14.98 ± 0.02 (15.01) | 16.29 ± 0.50 | 3.80 ± 0.06      |
| + L18W-PGLa/MG2a  | 63.29 ± 0.06   | 38.31 ± 0.03 | 14.68 ± 0.01 (14.93) | 15.98 ± 0.43 | 4.45 ± 0.04      |

*Values in brackets are derived from all-atom MD simulations.*
for the presently studied magainins as function of peptide concentration resulting from this analysis. In general, we find that the area expansion decreases with peptide concentration and levels off at \(-0.22\,\text{Å}^2/\text{K}\) independent of the specific structure of the added peptides, including their mixture. This shows that membranes are increasingly less able to expand laterally with increasing temperature as a result of the peptide-induced membrane perturbation.

**All-atom and coarse-grained simulations**

To gain additional insights on the molecular level, we performed MD simulations of both peptides within POPE/POPG (3:1 mol/mol) bilayers. Fig. S12 compares the form factors derived from all-atom MD simulations to experimental data. The remarkable agreement in particular for \(q > 0.15\,\text{Å}^{-1}\) provides a valuable validation of the results presented below.

**Individual peptides**

Firstly, we performed all-atom simulations with peptide monomers in which each membrane leaflet contained a single peptide to ensure membrane and system symmetry. Throughout the simulations, both peptides retained their mostly \(\alpha\)-helical conformation and remained oriented parallel with respect to the membrane plane (Fig. 4). The exception was MG2a, which showed a partial loss of helicity at the C-terminus in one leaflet. Fig. 4 shows snapshots from the end of the 500-ns-long simulations together with the depth of peptide insertion. L18W-PGLa was found to be slightly deeper in the headgroup region compared to MG2, which is consistent with our SANS/SAXS analysis.

We observed a local membrane modification in the vicinity of the peptides, in agreement with our scattering data analysis. Lipids changed their tilt and conformation to fill the hydrophobic void below the inserted peptide, as shown previously (51). To quantify this effect, we calculated the density distribution of the methyl groups of the lipid tails in the peptide aligned trajectories (Fig. 5, A and B). The methyl density was locally increased below the peptides, filling the available space between peptide side chains. The consequent effects on hydrocarbon chain packing are observed in the order parameter profiles’ dependence as a function of the distance from the peptides (Fig. 6, A and B; see also Figs. S13 and S14). In general, our analysis showed that palmitoyl chains are mostly affected at intermediate segments, whereas oleoyl hydrocarbons experience most significant changes toward the hydrocarbon tails. Moreover, MG2a appears to induce a slight increase of order close to the POPOP glycerol backbone of the unsaturated and a more pronounced disordering of its saturated hydrocarbon chain, respectively. Overall, the effects on hydrocarbon packing lead to membrane thinning, in good agreement with our scattering data analysis (see Table 1). The changes in density distributions for individual groups of membrane are depicted in Figs. S15 and S16.

**Effects of peptide dimers**

MG2a and PGLa peptides were previously shown to prefer parallel dimers using a coarse-grained model (52), which also agrees with experimental findings (7,9). We verified this tendency also for our bacterial membrane mimic. Because of the low number of dimer formation events at low peptide concentrations within the length of our simulations (20 μs), we had to increase P:L ratios from 1:42 to 1:21 to facilitate the analysis. We note, however, that this did not lead to the formation of a transmembrane pore, as observed in simulations using dilauroylphosphatidylcholine bilayers (52). This is due to the larger membrane thickness and tighter interfacial lipid packing of POPE/POPG (3:1 mol/mol) bilayers (5) and allowed us to derive the dimerization behavior even at elevated peptide levels. MG2a/L18W-PGLa mixtures, starting from random configuration, showed the strongest preference for dimerization, followed by MG2a and L18W-PGLa (Fig. 7). MG2a/ L18W-PGLa mixtures mainly formed parallel heterodimers, whereas MG2a and L18W-PGLa preferentially formed anti-parallel homodimers. Additional independent 40-μs-long coarse-grained simulations starting with peptides preformed in various heterodimer configurations corroborated the higher stability of parallel heterodimers as compared to other peptide-peptide alignments (Fig. S20). More insight into the stability of all dimer configurations could be obtained, in principle, from free-energy calculations; however, such calculations are very computationally demanding and are beyond the scope of this manuscript.

Because the peptide-peptide interactions might be overestimated in coarse-grained simulations (53), we performed additional all-atom simulations starting from preformed
dimers. In agreement with our coarse-grained simulations, we found that parallel heterodimers were stable on the simulated timescale.

Based on the preference of MG2a/L18W-PGLa mixtures to form parallel MG2a/L18W-PGLa heterodimers, we investigated the effect of this heterodimer on membrane structure using all-atom simulations of parallel heterodimers in both membrane leaflets (see Fig. S17 for final snapshot of the 500 ns trajectory). In agreement with experimental data, heterodimers lead to increased membrane thinning (Table 1) and significantly pronounced hydrocarbon chain packing defects (Fig. 6C) as compared to the individual peptides. This supports our above notion that the experimentally observed enhanced membrane perturbation is a result of L18W-PGLa/MG2a dimer formation at low peptide concentrations.

In the next step, we interrogated our all-atom simulations for the effect of dimerization on peptide location within bilayers. Compared to MG2a and L18W-PGLa (Fig. 4C), L18W-PGLa/MG2a heterodimers inserted more shallowly into the membrane (Fig. S18).

An analysis of our coarse-grained MARTINI simulations at both peptide concentrations showed that the trans-bilayer position of the heterodimers was within uncertainty almost equal to that of MG2a (Fig. 8; Fig. S19). To see whether the dynamics of dimer formation might affect their insertion depth, we artificially kept the peptides in parallel heterodimer configuration in an additional simulation (for details, see the Materials and Methods). This simulation decreased the uncertainty of the peptide position significantly but gave the overall same result for the dimer position as the unconstrained coarse-grained simulations, i.e., dimers are located closely to the phosphate group. The main difference compared to all-atom simulations is the slightly less deep insertion of MG2a, which could be due to force field issues. However, on an absolute scale, these differences are only minor. Moreover, results of both simulation models even agree considering the positional uncertainty of the peptides (Fig. 8; Figs. S18 and S19). Regarding experimental data (Table 1), we find an overall reasonable agreement with respect to the peptide position. Specifically, there is broad consensus between the different experimental and computational techniques applied here that L18W-PGLa positions itself further out from the bilayer center when dimerizing with MG2a.

**FIGURE 4** Membrane positioning of MG2a and L18W-PGLa peptides. Left side shows the snapshots from the end of 500-ns-long MD simulations of (A) MG2a and (B) L18W-PGLa adsorbed at the membrane surface. (C) shows the peptide positions in both leaflets averaged over trajectories. Snapshot color coding: lipid phosphate atoms are shown as orange spheres. Solvent is represented by a blue-shaded area, and lipid tails are not shown for clarity. The peptide secondary structure is shown in a cartoon representation, colored by residue type (nonpolar: gray; polar: green; acidic: red; and basic: blue). To see this figure in color, go online.

**FIGURE 5** Averaged densities of terminal hydrocarbon CH3 groups (cyan surfaces) after insertion of MG2a (A) and L18W-PGLa (B) in POPE/POPG bilayers. To see this figure in color, go online.
FIGURE 6 Changes of the lipid tail order parameters as a function of distance from the peptide (A) MG2a, (B) L18W-PGLa, and (C) MG2a + L18W-PGLa. To see this figure in color, go online.
We combined SAXS/SANS experiments with all-atom and coarse-grained MD simulations to interrogate mutual interactions between L18W-PGLa, MG2a, and fully hydrated POPE/POPG (3:1 mol/mol) bilayers. The focus of this work is on low peptide concentrations, i.e., at which equimolar mixtures of both peptides do not permeabilize POPE/POPG bilayers synergistically (5). Yet, we demonstrate that the peptide mixture leads to significantly enhanced membrane perturbations even at low concentrations.

We developed an SDP analysis for peptide-containing lipid membranes, which allowed us to determine the peptide position in the bilayer with high accuracy at low peptide concentrations by a simultaneous statistical analysis of four differently contrasted SAXS/SANS experiments. The observables were in excellent agreement with our MD simulations, enabling additional insight on lipid-peptide and peptide-peptide interactions. Note that previous reports on similar systems using solid-state $^{15}$N-NMR were not sensitive to the penetration depth of peptides within the headgroups. Both our experimental and simulation data consistently showed that MG2a inserts slightly less into the bilayer than L18W-PGLa (Fig. 4C; Table 1). This can be understood in terms of the larger hydrophilic surface of MG2a compared to L18W-PGLa because of the larger number of polar residues (5).

In agreement with $^{15}$N-NMR data on POPE/POPG (3:1 mol/mol) (13) and POPE-enriched bilayers (11), we found that both peptides adopt a surface-aligned topology, even in the case of equimolar L18W-PGLa/MG2a mixtures. The surface-aligned topology caused significant disorder of hydrocarbon chain packing, which was most pronounced in the vicinity of the peptides. In particular, we observed that the methyl termini of the hydrocarbons filled large fractions of the void below the peptide (Fig. 5), consistent with previous reports on dimple formation and membrane thinning (54). Membrane thinning was observed for both peptides (Table 1), consistent with a surface-aligned topology. The pronounced membrane thinning for equimolar mixtures of the two peptides provides indirect evidence for dimer formation. This is further supported by our experimental form factors, which were determined from a statistical data analysis and clearly show that the effects of the individual peptides cannot be simply combined to yield the membrane structure in the presence of both peptides (Fig. S5).

Hence, PGLa and MG2a seem to form dimers at concentrations much lower than reported previously (7,9,52).

Using all-atom MD simulations, we derived order parameter profiles as a function of distance from the peptides. In agreement with a previous NMR study (13) both peptides were found to perturb the saturated hydrocarbons of surrounding lipids (Fig. 6). Our results suggest that MG2a is more effective than L18W-PGLa in doing so, which correlates with the further outward location of MG2a within the membrane and its increased bulkiness (molecular volumes: $V_{L18W-PGLa} = 4927.8 \ \text{Å}^3$ and $V_{MG2a} = 5748.0 \ \text{Å}^3$). MD simulations additionally allowed us to probe order parameter profiles of unsaturated hydrocarbons. Interestingly, we found that MG2a increased the order of the oleoyl chain close to the glycerol backbone of POPG only, whereas a general decrease of this chain’s order parameter was observed for POPE at all segments for both peptides (Fig. 6, A and B; Figs. S13 and S14). This indicates specific
interactions of MG2a with POPG, which are located around the positively charged amino acids as revealed a detailed analysis of MD data (Fig. S21). Finally, assuming heterodimers, all effects observed with respect to the order of hydrocarbon chains were significantly more pronounced as compared to single peptides (Fig. 6 C), in agreement with our experimentally observed changes of membrane structure. This is also consistent with previous reports (see, e.g., (55)), showing that the formation of dimers amplifies the perturbation of the lipid tails in the vicinity of peptide dimer as compared to monomers.

Our simulations indicated that homodimers formed preferentially in antiparallel configuration, whereas L18W-PGLa/MG2a formed mainly parallel heterodimers (Fig. 7), which is in agreement with a previous study (52). Interestingly, heterodimer formation has been reported for significantly different lipid bilayers. Thus, dimerization of PGLa and MG2a does not appear to be highly specific to lipid composition, although we cannot comment on the onset of dimerization in other lipid bilayers from our study. The analysis of enthalpic interactions between different amino acids showed that dimer formation is stabilized by salt bridges between MG2a-Glu19 and Lys12 or Lys15 residues of L18W-PGLa (Fig. S22), in agreement with (52). Moreover, there are significant hydrophobic interactions between the peptides. For example, Ulmschneider et al. reported a stabilization of PGLa homodimers by Gly-Ala interactions at high P:L ratios (56). It appears, however, that the sum of all these interactions leads to a preferential formation of L18W-PGLa/MG2a heterodimers as compared to L18W-PGLa/L18W-PGLa or MG2a/MG2a homodimers, which is consistent with previous observations (6).

Most interestingly, peptide dimerization affects the penetration depth of the surface-aligned peptides. In particular, L18W-PGLa moves somewhat further out in the headgroup region when it associates with MG2a (Figs. 8, S18, and S19; Table 1). This leads to a larger void created in the membrane interior just below the peptides, causing an increased change of membrane structure. Hence, the ability to form a dimer appears to be important for the membrane perturbation efficacy of the studied peptides.

**CONCLUSION**

Our study suggests that an early onset of the formation of peptide dimers is the key event to the enhanced activity of L18W-PGLa/MG2a mixtures. Previously, we speculated that L18W-PGLa causes a deeper insertion of MG2a into the bilayer (5). Indeed, this work shows the opposite, i.e., that L18W-PGLa moves further out from the membrane center when forming a heterodimer with MG2a. These heterodimers perturb membranes significantly more than the sum of the effects induced by individual (noninteracting) peptides. Apparently, this “synergistic” dimerization is not sufficient to allow enhanced leakage of dyes (although smaller polar molecules might already permeate the bilayer) (5). For membrane leakage to occur, higher peptide concentrations are required. The corresponding membrane restructuring effects will be described in the subsequent study.

**SUPPORTING MATERIAL**

Supporting Material can be found online at https://doi.org/10.1016/j.bpj.2019.10.022.

**AUTHOR CONTRIBUTIONS**

M.P. performed the experimental research, analyzed the data, and wrote the article. I.K. carried out all simulations, analyzed the data, and wrote the article. M.-S.A. performed SANS experiments. R.V., K.L., and G.P. designed the research and wrote the article.

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**SUPPORTING CITATIONS**

Reference (57) appears in the Supporting Material.

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