

# Selective Interaction of Colistin with Lipid Model Membranes

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ABSTRACT Although colistin's clinical use is limited due to its nephrotoxicity, colistin is considered to be an antibiotic of last resort because it is used to treat patients infected with multidrug-resistant bacteria. In an effort to provide molecular details about colistin's ability to kill Gram-negative (G(-)) but not Gram-positive (G(+)) bacteria, we investigated the biophysics of the interaction between colistin and lipid mixtures mimicking the cytoplasmic membrane of G(+), G(-) bacteria as well as eukaryotic cells. Two different models of the G(-) outer membrane (OM) were assayed: lipid A with two deoxy-manno-octulosonyl sugar residues, and Escherichia coli lipopolysaccharide mixed with dilaurylphosphatidylglycerol. We used circular dichroism and x-ray diffuse scattering at low and wide angle in stacked multilayered samples, and neutron reflectivity of single, tethered bilayers mixed with collistin. We found no differences in secondary structure when collistin was bound to G(-) versus G(+) membrane mimics. ruling out a protein conformational change as the cause of this difference. However, bending modulus  $K_{C}$  perturbation was quite irregular for the G(-) inner membrane, where colistin produced a softening of the membranes at an intermediate lipid/peptide molar ratio but stiffening at lower and higher peptide concentrations, whereas in G(+) and eukaryotic mimics there was only a slight softening. Acyl chain order in G(-) was perturbed similarly to  $K_C$ . In G(+), there was only a slight softening and disordering effect, whereas in OM mimics, there was a slight stiffening and ordering of both membranes with increasing colistin. X-ray and neutron reflectivity structural results reveal colistin partitions deepest to reach the hydrocarbon interior in G(-) membranes, but remains in the headgroup region in G(+), OM, and eukaryotic mimics. It is possible that domain formation is responsible for the erratic response of G(-) inner membranes to colistin and for its deeper penetration, which could increase membrane permeability.

## INTRODUCTION

Colistin (i.e., polymyxin E) was first isolated by Koyama et al. (1) from the broth of *Bacillus polymyxa* in 1949; it is a linear trilipopeptide linked to a cyclic heptapeptide that is produced by nonribosomal peptide synthetase systems in Gram-positive (G(+)) bacteria (2). The fatty acid tail can consist of seven, eight, or nine carbons (3). It has a narrow antibacterial spectrum, mainly against Gram-negative bacteria (G(-)) (4,5), but not G(+) bacteria. Clinical use of colistin decreased in the 1970s due to nephrotoxicity and neurotoxicity after intravenous administration. However, the world is now facing a growing threat from bacteria that are resistant to all available antibiotics (6,7), and colistin has reemerged as an antibiotic of last resort (3), and is in use to treat cystic fibrosis patients. The

\*Correspondence: stn@cmu.edu Editor: Georg Pabst. https://doi.org/10.1016/j.bpj.2017.12.027 © 2017 Biophysical Society. incidence of resistance to colistin is relatively low (8), but resistance in Gram-negative pathogens can emerge both in vitro (9,10) and clinically (11-14).

Colistin interacts with high affinity with the lipopolysaccharide (LPS) of the outer membrane (OM) of G(-) bacteria (15), which docks the positively charged (+5 e) antibiotic to the cell surface. A critical interaction of colistin with the lipid A component of LPS is suggested by the charge screening mechanism of bacterial resistance whereby negative phosphate charges are neutralized by the addition of aminoarabinose and ethanolamine residues (16). However, membrane blebbing (17) and electrochemical transmembrane potential dissipation in cells treated with colistin have also been reported in previous works (18), which suggests that inner membrane (IM) permeabilization leads to bacterial cell death. What is the mechanism of colistin's rapid, concentrationdependent bacterial killing with negligible postantibiotic effects (10)? Hancock and co-workers (19–23) have proposed a general model for several antimicrobial peptides called the



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"self-promoted uptake" model, where colistin's aggregation promotes its own uptake across the OM, and subsequent pore formation of the IM. An alternative proposed mechanism is the vesicle-vesicle contact pathway (24,25) where a colistin dimer can mediate the contacts between periplasmic leaflets of inner and outer membranes (24). A third possible mechanism is a generalized mechanism for bactericidal agents, in which an oxidative burst produces a reactive hydroxyl radical (•OH) that can induce rapid cell death (26).

This work aims to delineate the role of lipids in the interactions between colistin and bacterial lipid membrane mimics in an effort to understand the molecular details of colistin's bactericidal mechanism. It is of interest to determine if there are structural or elastic properties that differ between G(+) and G(-) lipid membrane mimics that could be responsible for colistin's preferential killing of G(-) bacteria. As a first approach, we studied the secondary structure of the peptide by means of circular dichroism in the absence and presence of lipid membranes mimicking the outer leaflet (OM) and IM of G(-), cytoplasmic membrane of G(+) bacteria, and eukaryotic cells. With the same membrane mimics, by means of x-ray diffuse scattering we measured both structure (membrane thickness, area/lipid, and peptide position) and the elastic parameter bending modulus  $(K_C)$ , which yields information about membrane softening. In addition, an order parameter that indicates lipid acyl chain ordering is obtained  $(S_{xray})$ . Finally, the third biophysical method, neutron scattering, was used for confirming colistin's location in the membrane mimics. Because our experimental x-ray system is symmetric, we cannot probe the vesiclevesicle contact model, and we are also not studying oxidized lipids, so we cannot comment on these other two models for bactericidal killing.

# MATERIALS AND METHODS

#### Reagents

The synthetic lyophilized lipids 1-palmitoyl-2-oleoyl-*sn*-glycero-3phosphoethanolamine (POPE), 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phospho-(1'rac-glycerol) sodium salt (POPG), 1', 3'-bis[1,2-dioleoyl*sn*-glycero-3-phospho]-*sn*-glycerol sodium salt (TOCL, i.e., cardiolipin), 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC), 1,2-dioleoyl-3-trimeathylammonium-propane chloride salt (DOTAP), di[3deoxy-D-manno-octulosonyl]-lipid A ammonium salt (KDO2), and 1,2-dioleoyl-*sn*-glycero-3-[phospho-*rac*-(3-lysyl(1-glycerol))] (chloride salt) (Lysyl PG) were purchased from Avanti Polar Lipids (Alabaster, AL) and used as received. Cholesterol was from Nu-Chek-Prep (Waterville, MN). HPLC grade organic solvents, LPS from *Escherichia coli* 0111:B4, and colistin sulfate salt were purchased from Sigma-Aldrich (St. Louis, MO) and used as received.

#### Sample preparation

Membrane mimics were prepared by first dissolving lyophilized lipids in chloroform/methanol (8:2, v/v), or LPS in chloroform/methanol/H<sub>2</sub>O (2:1:1, v/v/v). Lipid stock solutions were mixed to create lipid mixtures in

molar ratios mimicking bacterial membranes: G(-) IM, POPE/POPG/ TOCL (7:2:1, molar ratio); G(+) membrane, POPG/DOTAP/POPE/TOCL (6:1.5:1.5:1); and eukaryotic membrane, POPC/POPE/cholesterol (5:1:1.8). For the outer membrane of G(-) bacteria, two different mimics were used: lipid A with two deoxy-manno-octulosonyl sugar residues of the inner core (KDO2), and a mixture of LPS/DLPG (1:3). Stock solutions of colistin (MW = 1400) were prepared in Milli-Q water (Millipore, Billerica, MA).

Multilamellar stacked samples for x-ray scattering were prepared by mixing 4 mg of the different lipid mixtures plus colistin into a glass test tube in various mole ratios from 750:1 to 50:1, lipid/peptide. Solvents were removed by evaporation under vacuum and samples were redissolved in appropriate HPLC-grade solvents for spreading: G(-), G(+), chloroform/trifluoroethanol (TFE) 7:3 (v/v); eukaryotic, chloroform/TFE 1:1 (v/v); LPS model, chloroform/TFE/H<sub>2</sub>O (5:5:1); and KDO2, chloroform/ methanol/H<sub>2</sub>O (90:10:1). These mixtures were plated onto silicon wafers (15 × 30 × 1 mm) via the rock-and-roll method (27) to produce stacks of ~1800 well-aligned bilayers. After solvent removal under vacuum for 2 h, hydration occurred through the vapor in a thick-walled x-ray hydration chamber (28).

Samples for circular dichroism (CD) spectroscopy were prepared by spreading a thin film of 0.7 mg of the same bacterial mimic mixtures as for x-ray with/without 0.3 mg colistin onto the inner wall of a 1-cm quartz cuvette. After solvent removal under vacuum, CD samples were fully hydrated at room temperature overnight with 100  $\mu$ L Milli-Q water in the bottom of the sealed cuvette.

Samples for densimetry were prepared as in (29). A quantity of 10–50 mg dried lipid mixture was mixed with  $\sim$ 1.2 mL water. This mixture was hydrated by temperature cycling three times from 60 to 0°C with vortexing to produce multilamellar vesicles.

Samples for neutron scattering were prepared by adding colistin to 8 mg lipid mixtures in a 50:1 lipid/peptide molar ratio. Organic solvent was removed by evaporation and samples were rehydrated in a 2 M NaCl aqueous solution to a final concentration of 5–6 mg/mL and bath-sonicated for 60–90 min until clarity. The self-assembled monolayers of HC18 tethers were formed on 3" diameter silicon wafers (30) and sparsely tethered bilayer lipid membranes were formed by exposing the self-assembled monolayer to the vesicle suspension for 60 min in a National Institute of Standards and Technology (NIST) reflectivity flow cell. This was followed by a rinse with 40 mL deionized water (31).

#### Methods

Low-angle x-ray scattering. Low-angle x-ray scattering (LAXS) data from oriented, fully hydrated samples were obtained at the G1 line at the Cornell High Energy Synchrotron Source (CHESS, Ithaca, NY) with previously described methods (32–34) on two separate trips using x-ray wavelengths 1.108 and 1.096 Å and sample-to-detector (*S*)-distances of 387.2 and 396.6 mm. In addition, a laboratory x-ray source RUH3R Rotating Anode X-Ray Generator (Rigaku, Tokyo, Japan) with a FOX 2D Focusing Collimator (Xenocs, Sassenage, France) and a Mercury CCD Detector (Rigaku) were used with an x-ray wavelength of 1.5418 Å and S-distance of 280.6 mm. Full hydration is judged by no further increase in lamellar D-spacing over time. Measurements were carried out in the fluid phase typically at 37°C, except for KDO2 samples, which were studied at 55°C due to the high melting temperature of the lipid. Details about determination of  $K_C$ from diffuse LAXS images and electron density profiles are given in the Supporting Material.

*Wide-angle x-ray scattering.* Wide-angle x-ray scattering (WAXS) was obtained at CHESS (*S*-distances = 163.4 and 179.3 mm) and at Carnegie Mellon University (Pittsburgh, PA) (*S*-distance = 125.7 mm) as described in (29,35). Hydrocarbon chain order parameters were estimated by measuring the angular dependence of the interchain WAXS signal according to the model developed by Mills et al. (35), which is also based upon liquid crystal theory. More details of the WAXS analysis are given in the Supporting Material.

*Neutron reflectivity*. Neutron reflectivity (NR) measurements were performed at the NGD-MAGIK reflectometer at the NIST Center for Neutron Research (Gaithersburg, MD) over a momentum transfer range 0-0.25 Å<sup>-1</sup>. More details about NR are given in the Supporting Material.

*CD spectroscopy.* CD spectra in the 200–240-nm range were collected with a model No. 715 Spectropolarimeter (Jasco, Oklahoma City, OK) by accumulating 15 or 20 spectra at 37°C, at 100 nm/min with a step resolution of 1 nm. Samples were performed in duplicate or triplicate. More details of the CD spectroscopy are given in the Supporting Material.

Densimetry. Approximately one milliliter of multilamellar vesicles were loaded into the model No. 5000M Densimeter (Anton-Paar, Graz, Austria). Density measurements were recorded as in (36), by first measuring Milli-Q water at 37°C and then measuring the sample density at 37°C. The sample was withdrawn and remeasured several times to ensure reproducibility. The molecular volume was calculated from density using Avogadro's number and the combined molecular weight of the lipids assuming additivity. Colistin's volume was measured separately as an aqueous solution at 37°C (7.9 mg/mL).

#### RESULTS

#### CD

Colistin adopted a secondary structure in water that contained primarily  $\beta$ -sheet, random coil, and  $\beta$ -turn (see Table 1). When associated with lipids in thin films, however,  $\beta$ -turn and  $\beta$ -sheet structure were reduced, and a signal resembling  $\alpha$ -helix could be fitted. However, it must be recalled that in peptides, some turns can show the same negative band at 222 nm as a result of n- $\pi$ \* transitions (37,38). What is striking is that colistin's secondary structure is quite similar in all of the lipid models, with slightly higher  $\alpha$ -helix-type signal in LPS and eukaryotic membranes. In particular, there is little difference between colistin in G(-) and G(+) membrane mimics. Light microscopy pictures of the thin film samples in the dried state that were then hydrated for CD are shown in Fig. S1. CD ellipticity data that produced the results in Table 1 can be found in Fig. S2.

#### **Diffuse scattering**

Fig. S4 shows typical LAXS data from oriented, fully hydrated stacks of membrane mimics containing colistin, 100:1 lipid/colistin. The results of the LAXS fitting are shown in Figs. 1, *A* and *B*, 2, and 3.

Fig. S7 shows WAXS data obtained from the same sample concentrations as in Fig. S4. The results of the WAXS fitting are shown in Fig. 1, *C* and *D*.

TABLE 1 CD Results of Colistin in Water and Membrane Mimics

Colistin Sample	α-Helix- Type %	β-Sheet %	β-Turn %	Random Coil %	$R^2$
Water	$0 \pm 1$	48 ± 6	16 ± 5	36 ± 8	0.88
G(-)	$24 \pm 6$	$18 \pm 8$	$0 \pm 1$	$59 \pm 7$	0.99
G(+)	$17 \pm 3$	$17 \pm 4$	$2 \pm 1$	$64 \pm 2$	0.99
LPS	$31 \pm 11$	$3 \pm 3$	$0 \pm 1$	$67 \pm 3$	0.99
Eukaryotic	$30 \pm 10$	$9 \pm 8$	$0 \pm 1$	$62 \pm 12$	0.99

Percent composition was obtained by normalizing to 100% the unitless coefficients in the linear least squares fit of the structural motifs to the data.  $R^2$  values indicate goodness of fit, with 1 indicating a perfect fit to the data. Bending moduli and  $S_{xray}$  order parameters

Fig. 1 shows the results of fitting the LAXS (Fig. 1, A and B) and WAXS (Fig. 1, C and D) diffuse data for the five lipid membrane mimics containing colistin. Not surprisingly,  $K_C$  is highest (stiffest membrane) for the eukaryotic membrane mimic, in which cholesterol interacts favorably with the saturated palmitoyl chain (39) in POPC and POPE (Fig. 1 A). Colistin caused only a slight softening of the eukaryotic mimic with increasing concentration. G(+) membrane mimics had a much lower  $K_C$ , indicating a more bendable membrane, which was also only slightly softened by colistin. G(-), on the other hand, showed a complex behavior of  $K_C$  with colistin concentration. At low (500:1) and high peptide concentration (>100:1 lipid/peptide molar ratio), colistin induced stiffening, but at intermediate ratios  $(\sim 200:1)$ , a significant membrane softening was obtained (Fig. 1 A). For studying the interaction of colistin with the outer membrane of G(-) bacteria, a model mixture consisting of LPS/DLPG (1:3 molar ratio) was employed, because multilayered stacks of pure LPS showed a lamellar x-ray pattern with D-spacing of ~290 Å but no diffuse signal, indicating nonfluctuating bilayers and impairing the analysis of mechanical and structural properties of the membrane within the theoretical framework of liquid crystals. The  $K_C$  values in Fig. 1 B for LPS/DLPG model were lower than those for the G(+) membrane mimic, indicating that the model used for OM is a relatively soft membrane, whereas DLPG alone (*black triangle*) had a higher  $K_C$ , indicating that the additional sugar residues on LPS cause a softening. KDO2 stacked bilayers in fluid phase at 55°C had a higher  $K_C$  than LPS/DLPG at 37°C and showed stiffening with increasing colistin content (Fig. 1 B). In Fig. 1, C and D, the complementary  $S_{xray}$  results for the  $K_C$  results in Fig. 1, A and B are shown.  $S_{xray}$ , which indicates acyl chain order, paralleled membrane bending in all of the membrane mimics. As shown in Fig. 1 C, eukaryotic membrane, containing cholesterol, had the most ordered chains, and colistin only slightly disordered them. G(+) had the most disordered chains and colistin slightly disordered them. G(-) showed the same irregular behavior for  $S_{xrav}$  as for  $K_C$ , first increasing chain order at 500:1 lipid/peptide molar ratio, then decreasing chain order at 200:1, then increasing chain order to a maximum at 100:1. In the OM mimics in Fig. 1 D, the chain order increased slightly with increasing colistin concentration, similar to the slight membrane stiffening shown in Fig. 1 B. Similar to its  $K_C$  effect, KDO2 had a higher chain order compared to LPS model. LPS model also has more disordered chains than DLPG, suggesting a chain disordering effect of the sugar residues.

#### Structural results

Fig. 2 shows the electron density profiles (EDPs) of the five membrane mimics containing the highest concentration in our study of colistin for each mimic. The measured volumes



FIGURE 1 Elasticity ( $K_C$ ) results for (A) eukaryotic, G(-) and G(+) membrane mimics, (B) KDO2 and LPS model, and DLPG control.  $S_{xray}$  order parameter results for (C) eukaryotic, G(-) and G(+) membrane mimics, (D) KDO2 and LPS model, and DLPG control. To see this figure in color, go online.

required for determining the EDPs appear in Table S1. Colistin (solid magenta, Gaussian) showed a deeper partitioning into the G(-) IM membrane mimics (Figs. 2 C and 3 C) when compared to both G(+) and eukaryotic mimics at the different lipid/peptide mole ratios assayed. In G(-)IM, colistin is located in the hydrocarbon region just within the interfacial region, 11-14 Å from the bilayer center, at all concentrations. In eukaryotic and G(+) membrane, colistin locates in the headgroup region, just within  $D_{\rm HH}/2$ . Colistin in G(-) mimics locates in the headgroup region in KDO2 (Fig. 2 A) and outside of the headgroup maximum  $(D_{\rm HH}/2)$  in the mixture LPS/DLPG (1:3) (Fig. 2 B). The EDP of the LPS/DLPG mixture was calculated taking into account the electrons from the Re variant of the LPS molecule (lipid A plus the inner sugar core, consisting of three residues of deoxy-manno-octulosonyl and 2 phospho-heptose), whereas the remainder sugar moieties in the core and O-antigen regions were included in the water electron density.

Fig. 3 summarizes the structural results for all five membrane mimics with increasing colistin concentration.  $A_L$  is the area/unit cell, which includes the lipid and the colistin, but not the cholesterol (for eukaryotic membrane). In G(-) IM,  $A_L$  (black lines) decreased at 500:1 lipid/peptide molar ratio but increased at 200:1. These changes are in the expected direction according to the  $S_{xray}$  results in Fig. 1 *C*, because increasing chain order decreases  $A_L$ , as the chains straighten. At peptide concentrations higher than 100:1,  $A_L$  showed no significant differences when compared to control. The overall bilayer thickness ( $D_{HH}/2$ , *red lines*) did not change appreciably. In G(+) membranes,  $A_L$  first decreases slightly then increases with colistin concentration, whereas in eukaryotic membranes, there is an increase in  $A_L$  with colistin concentration.  $A_L$  is fairly constant as colistin is added to KDO2 and LPS model, as is the bilayer thickness,  $D_{\rm HH}/2$ . Table S4 compares our KDO2 control area with literature values.

### NR

Fig. 4 shows the NR results as component volume occupancy versus distance from the gold substrate. At this concentration of lipid/peptide (50:1), colistin (*red line*) enters more deeply into the G(-) membrane (Fig. 4 *A*) than into the G(+) membrane (Fig. 4 *B*). However, compared to the x-ray result, there is considerable colistin outside of the outer leaflet headgroup Gaussian (*cyan*) for both G(-) and G(+) membrane mimics. This could be due to the different sample preparation using vesicle fusion to form a single tethered bilayer, compared to the stacked system of 1800 bilayers for x-ray. The significance of the NR result is that there is a deeper penetration of colistin into the hydrocarbon core in G(-) membrane mimics, similar to the x-ray result. Additional details of the NR results are shown in the Supporting Material.

### DISCUSSION

# Colistin secondary structure is similar in different membrane mimics

A shape analysis procedure (38) for fitting the experimental CD spectra was undertaken for estimating the secondary



FIGURE 2 EDPs for (*A*) KDO2, 50:1 lipid/peptide molar ratio; (*B*) LPS model, 50:1; (*C*) IM, 75:1; (*D*) Eukaryotic, 50:1; (*E*) G(+), 100:1. Component groups are headgroup HG1 (*red*), headgroup HG2 (*violet*), methylene CH2 region (*blue*), methyl CH3 region (*green*), colistin (*solid magenta*), water (*cyan*), and total (*black*). To see this figure in color, go online.

structure of colistin, using a set of spectra of typical motifs obtained from standard proteins/polypeptides (40). Although these reference spectra correspond to proteins formed by the naturally occurring L isomer of amino acids, we employed them for this colistin analysis, whose primary structure is formed of nine L-amino acids plus one D-amino acid. This method was appropriate because several previous reports indicated that CD is a sensitive measurement of the secondary structural changes caused by single or double D-amino acid substitution in short peptides (41-43). On the other hand, complete inverse CD structure is obtained in an all-D model  $\alpha$ -helixes (44,45). Our study indicates that, in agreement with previous NMR results (46,47), the colistin backbone in the aqueous phase has a significant degree of freedom, with a high random coil or disordered structure plus  $\beta$ -turn and  $\beta$ -sheet. When bound to lipids, both  $\beta$ -turn and  $\beta$ -sheet content decreased whereas disordered structure increased, which agrees with the weakening of the  $\beta$ -turn NMR signal observed previously (46). However, the  $\alpha$ -helix-type feature was also found, which could be ascribed to distorted turns, as previously observed for other cyclic peptides and D-amino acid-containing peptides (43).

The further result that there were no significant differences in secondary structure of colistin when bound to the different membrane mimics, suggests that a change in colistin secondary structure cannot be the reason for differences in bactericidal mechanism in G(-) versus G(+) bacteria.

## Colistin's effects on G(–) IM membrane mechanical properties and order parameters may involve lateral heterogeneity

Contrary to the CD results, dramatic changes were observed in the elastic behavior of colistin interacting with G(-) IM mimics compared to G(+) mimics. As shown in Fig. 1 A, colistin caused alternating stiffening, softening and stiffening, as its concentration increased in G(-) IM mimics. This erratic behavior was mirrored by the  $S_{xray}$  order parameter (Fig. 1 C), where colistin alternately ordered, disordered and ordered the acyl chains at the same concentrations as for  $K_{C}$ . These changes are unlike any seen in our previous investigations: increasing concentration of HIV-1 fusion peptide (48) or alamethicin (49) decreased the bending modulus  $K_C$  in DOPC and diC22:1PC in an exponential fashion, and increasing HIV-1 matrix<sub>31</sub> peptide caused a gradual increase in  $K_C$  and  $S_{xray}$  then a gradual decrease in both in PS-containing membranes (29). The fact that this sharply irregular behavior occurred only in G(-) IM mimics may offer a clue to colistin's bactericidal mechanism in G(-)bacteria. Compared to G(+) mimics, which displayed only a slight softening and disordering of lipid chains, G(-) IM mimics were greatly perturbed, in both directions, by colistin.



FIGURE 3 Structural parameters, area/lipid  $A_L$  (black lines),  $D_{HH}/2$  (red lines) and colistin position (blue lines), as a function of increasing colistin mole fraction in five lipid membrane mimics: (A) KDO2, (B) LPS model, (C) G(-) IM, (D) Eukaryotic, and (E) G(+). Left axis shows  $A_L$  (black); right axes show  $D_{HH}/2$  (red) and peptide position (blue).  $D_{HH}/2$  and peptide position are in Å units from the bilayer center, and  $A_L$  is in units of Å<sup>2</sup>. To see this figure in color, go online.

The G(+) mimic used in this work consisted of the lipid mixture POPG/DOTAP/POPE/TOCL (6:1.5:1.5:1 molar ratio), as many Gram-positive bacteria, such as *Staphylococcus aureus*, contain a high amount of the negatively charged lipids PG and cardiolipin, but also a positively charged derivative of PG, lysyl-PG, in which a lysine residue is bonded to the PG headgroup (50). Due to the high amounts of lipids needed for preparing samples in a wide range of peptide concentrations, a more affordable lipid, DOTAP, was used instead of lysyl-PG, which is also positively charged. A control sample prepared with lysyl-PG showed no significant differences in  $A_L$  or  $K_C$  compared to mixtures prepared with DOTAP, indicating that the latter is a good surrogate for lysyl-PG. Unlike G(+) membranes, the inner membranes of G(-) bacteria are highly enriched in PE but also contain a significant amount of negatively charged lipids (51,52), so this mimic contained POPE/POPG/TOCL (7:2:1).

It has been suggested that cardiolipin-rich domains exist in G(-) membranes (53) and similar cationic antimicrobial peptides can perturb bilayer permeability by inducing phase separation in membranes containing PE and cardiolipin (54). Thus, we propose that colistin may induce membrane lateral heterogeneity by clustering anionic lipids due to Coulombic interactions in the presence of high content of PE, as in IM G(-) membranes but not in a low content of PE, as in G(+) membranes. Formation of lipid domains can lead to interfacial curvature stress in bilayers, due to a nonbilayer tendency that some of them have, like PE (55) or cardiolipin (56). A previous work found that the increase



FIGURE 4 Component volume occupancy versus distance from substrate. (A) G(-) IM mimic; (B) G(+) mimic. Lipid/peptide 50:1 molar ratio. Component groups are shown: water (*cyan*), colistin (*red*), headgroups (*olive*), hydrocarbons (*blue*), tether (*green*), and substrate (*gold*). To see this figure in color, go online.

in current due to permeability perturbation in asymmetric model membranes occurred above a threshold colistin concentration, which was also postulated to be a consequence of preferential binding of the peptide with negatively charged lipids and phase separation (57). The presence of lipid domains and interfacial curvature stress would explain transient membrane permeabilization along the defects at domain boundaries (54). In a recent molecular dynamics (MD) simulation (58), it was shown that the bending modulus was altered as a cationic AMP was added to a binary lipid membrane containing the zwitterionic DPPC and the negatively charged DPPS. The authors suggested that when cationic AMPs encounter bacterial membranes, domains with different material moduli are formed, which could lead to a destabilization at the boundaries of these domains. To investigate this possibility further, we have studied the  $K_C$  values of the individual components of G(-) IM, with and without 0.01 mol fraction colistin, with results shown in Table 2.  $K_C$  of POPG is reduced by about half when colistin is added, whereas  $K_C$  of TOCL is only slightly reduced. POPE with 0.01 colistin, or even 0.001 colistin, did not fluctuate, indicating that  $K_C$  has been greatly increased by colistin (i.e., membrane is stiffer). Thus, even though PE is the lipid at highest proportion in G(-) IM, the softening of the mixture at intermediate colistin mole fractions should be produced by a preferential interaction of colistin with PG or TOCL due to Coulombic forces, whereas the stiffening caused by colistin-PE interaction is evidenced at intermediate and higher concentrations. When these results are compared to Fig. 1 A, we might interpret the initial slight lowering of  $K_C$  as due to colistin binding to TOCL. As concentration increases, colistin could bind to POPE, thus increasing  $K_C$ . When collistin encounters POPG, a larger decrease in  $K_C$  is observed. At the highest concentrations, colistin interacts with the remaining POPE, causing a final increase in  $K_C$ . A cartoon summarizing this scenario is presented in Fig. 5.

A second reason for differences between G(+) and G(-) IM is that colistin reaches the deepest location in the hydrocarbon region in G(-) IM when compared to the other membrane mimics. The EDPs and summary of structural results show that colistin is located in the headgroup region at all concentrations for all mimics except for G(-) IM. NR supports this location, although the contrast between G(-) and G(+) is not as dramatic as in x-ray, presumably due to the different sample preparations. This deeper location of colistin in G(-) IM could cause a defect that would

TABLE 2 Bending Moduli,  $K_C$  Values, of Components of G(-) Inner Membranes

Colistin Mole Fraction	POPE	POPG	TOCL
0	$9.7 \pm 1.0$	$7.5 \pm 0.3$	$4.4 \pm 0.4$
0.01	ND	$3.7~\pm~0.4$	$3.3 \pm 1.0$

 $K_c$  units (×10<sup>-20</sup> J). ND, none detected.



FIGURE 5 Cartoon showing colistin interacting with individual lipid types in G(–) IM as its concentration increases. (A) By comparing to Fig. 1 A, colistin may first interact with TOCL, slightly lowering  $K_C$ . (B) Colistin may then interact with POPE, increasing  $K_C$ . (C) Colistin may then interact with POPG, decreasing  $K_C$  substantially. Finally, colistin may interact with the remaining POPE (data not shown), causing an increase in  $K_C$ . To see this figure in color, go online.

allow for a deeper penetration of water into the bilayer, leading to permeabilization and bacterial cell death.

# Colistin locates in the headgroup region in G(-) OM mimics

As for the G(-) OM mimics, we observe that the LPS model (LPS/DLPG 1:3) is softer and less ordered than DLPG or KDO2, indicating that the LPS component with its core and O-antigen sugar residues induces a softening and disordering effect. This contradicts the idea that the OM has low fluidity (59,60), but is in agreement with order parameters calculated by MD simulation that showed  $|S_{CD}|$  decreases with increasing sugar residues on lipid A (61). The hydrocarbon half-thickness for the two OM control mimics (EDPs not shown) are both thinner than usual ( $D_C = 12.2$  Å for KDO2 and 9.2 Å for LPS model), whereas the other control mimics have more normal thicknesses ( $D_C = 14.4$  Å for G(+), 14.8 Å for G(-), and 15.4 Å for eukary-otic). We find that colistin slightly orders both KDO2 and

the LPS model, contradicting the Fourier transform infrared spectroscopy results of (62), which showed a fluidizing effect of the related polymyxin B on KDO2. However, in that study, the smallest concentration of lipid/peptide was 8:1, a much larger amount of colistin than in our study. The small effect of colistin in our model OM systems agrees with previous results with polymyxin B, in which lower charge screening and permeabilization are observed in LPS- compared to PG-containing lipid vesicles (63).

Colistin's location in the headgroup of KDO2 and the LPS model suggests that the outer membrane destabilization observed in previous biophysical and microbiological works must occur through divalent cation displacement by colistin (18,64). Although our study did not add  $Ca^{2+}$  or  $Mg^{2+}$  as variables, our main conclusion, that colistin remains in the headgroup region of G(-) OM models, is consistent with the idea that it could displace divalent cations. Future experiments could explore the dependence of colistin's location on type of divalent cation. The divalent cations play an important role in the assembly of the LPS molecules in the outer membrane by screening the repulsive Coulombic forces between the phosphate residues at the lipid A and sugar inner core levels (59). It was previously reported that colistin binding to lipid A is inhibited by divalent cations and high ionic strength (64). By weakening the outer membrane, colistin could induce its self-promoted uptake toward the periplasmic space and reach its final target, the inner membrane of G(-)bacteria. A recent MD simulation confirmed our headgroup location of colistin in KDO2 and the LPS model (65).

## CONCLUSIONS

Using a structural and materials approach, this work attempted to clarify molecular steps in colistin's bactericidal mechanism. We found no difference in secondary structure of colistin in G(-) IM and G(+) membranes, thus ruling out a protein conformational change as the cause of colistin's ability to kill G(-) but not G(+) bacteria. However, dramatic differences between colistin's effect on G(-) IM versus G(+)membrane mimics were observed in the elasticity results; whereas G(-) IM mimics were softened at a critical lipid/ peptide molar ratio (200:1), they were stiffened above and below that ratio. G(+) membranes, on the other hand, were only slightly softened, a very small perturbation. We suggest that colistin induces domains in G(-) IM with different adjacent elasticity, which could lead to permeation through the domain boundaries. Chain ordering paralleled membrane elasticity to indicate that lipid acyl chains are significantly perturbed only in the case of G(-) IM. In addition, colistin located at a deepest position in the interior of the hydrocarbon region in G(-) membranes. Therefore, the elasticity, chain ordering, and peptide membrane location point to colistin's ability to enter into and perturb the G(-) inner membrane, which could lead to an increase in permeability. As for the G(-) OM, collistin remains in the headgroup region at all

ing ef-<br/>wer, inthe OM stiffness and chain order. Therefore, colistin is in<br/>position to displace divalent cations, leading to OM perturba-<br/>tion, and eventually, to its self-promoted uptake.study.

## SUPPORTING MATERIAL

Supporting Materials and Methods, ten figures, and four tables are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(17)35132-9.

the concentrations that we studied, and increases slightly

### **AUTHOR CONTRIBUTIONS**

F.G.D. and S.T.-N. designed the experimental research, performed experiments, analyzed data, and wrote the manuscript. I.P., K.A., and F.H. performed experiments and analyzed data. M.F.P. and Y.E. performed experiments.

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### SUPPORTING CITATIONS

References (66–79) appear in the Supporting Material.

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