ABSTRACT: Host defense or antimicrobial peptides hold promise for providing new pipelines of effective antimicrobial agents. Their activity quantified against model phospholipid membranes is fundamental to a detailed understanding of their structure−activity relationships. However, classical characterization assays often lack the ability to achieve this insight. Leveraging a highly parallelized microfluidic platform for trapping and studying thousands of giant unilamellar vesicles, we conducted quantitative long-term microscopy studies to monitor the membrane-disruptive activity of archetypal antimicrobial peptides with a high spatiotemporal resolution. We described the modes of action of these peptides via measurements of the disruption of the vesicle population under the conditions of continuous peptide dosing using a range of concentrations and related the observed modes to the molecular activity mechanisms of these peptides. The study offers an effective approach for characterizing membrane-targeting antimicrobial agents in a standardized manner and for assigning specific modes of action to the corresponding antimicrobial mechanisms.

The COVID-19 pandemic has refocused the world’s attention on the dramatic consequences of failing to contain infectious diseases. However, the "silent" pandemic of antimicrobial resistance (AMR) continues unabated and remains mostly out of the public eye. The spread of multidrug-resistant bacteria, combined with both economic and scientific challenges in drug development, has led to a situation, where common infections are becoming untreatable. There is a dire need to provide new pipelines of antimicrobial agents exhibiting mechanisms which may be less subject to acquired resistance.

Molecular agents disrupting microbial membranes are often advantageous over traditional antibiotics targeting individual intracellular processes in microbial cells. Their activity is not necessarily confined to a single target, does not require reaching the cytoplasm, but is not limited to membrane disruption either, allowing them to tackle a microbial cell as a whole. Such agents are of particular interest for treating Gram-negative bacteria whose double membranes continue being a formidable barrier for antibiotics with intracellular targets. Host defense or antimicrobial peptides (AMPs) are typical representatives of membrane-active antimicrobials. These peptides favor attack on bacterial cell surfaces. Due to their membrane activity, they avoid common resistance mechanisms (e.g., reduced influx or enhanced efflux) associated with drugs that affect intracellular targets and therefore show high efficacy against multidrug- and pandrug-resistant bacteria in vitro.

Yet, the translation of these agents and treatments into clinical use has not been efficient, suggesting a discordance between in vitro tests and clinical trials when attempting to balance out the efficacy, stability, and toxicity of these peptides. As the field has also recently noted, better and standardized characterization platforms are needed to facilitate a deeper and more systematic understanding of the antimicrobial mechanisms of these peptides. This will support more predictive structure–activity outcomes and hence facilitate the efficient clinical translation of AMPs.

When developing these tools, the terms "mechanism" and "mode of action" are often used interchangeably. In this study, we distinguish between the two in that "mechanism" refers to the dynamics governing interactions between peptides and phospholipids at the molecular level. This mechanistic understanding can be elucidated by employing techniques such as molecular dynamics (MD) simulations, oriented circular dichroism spectroscopy, and atomic force microscopy (AFM). By "mode of action", we mean the activity of
the peptides at the cell population level using either live cells or model membranes such as unilamellar vesicles. This may involve the kinetics of membrane disruption as determined by the distribution of leakage timepoints within a population of vesicles. With a molecular mechanism underpinning a mode of action, we chose to make this distinction explicitly. Peptide—membrane interactions are postulated to occur via the so-called “two-state model”; the surface state, during which peptides bind to membrane surfaces, and the insertion state when bound peptides insert into the membrane forming pores and channels. In this model, the point of transition between the two states is defined by reaching a threshold concentration of bound peptide after a certain lag period. The insertion state can be followed by pore expansion causing critical membrane disruption and eventually leading to cell lysis. The two-state model mechanism should manifest as a mode of action, in which all vesicles within a population are disrupted within a narrow band of lag times that is dependent on the dosage applied. Therefore, by studying the mode of action, we should then be able to discriminate between peptides, which subscribe to the two-state model and those that do not. For peptides which do not follow the two-state model, they fall into the “stochastic” mode of action, where the population of single vesicles is perturbed in a stochastic manner, manifested in peptide-induced disruption occurring in some vesicles, but not in others, or after random lag time periods independent of peptide concentration.

Leakage studies using reconstituted synthetic unilamellar vesicles provide a straightforward probe of the antimicrobial activity of membrane active AMPs. Typically, a fluorescent probe is encapsulated in the lumen of a phospholipid vesicle, and a stable (non-decaying) fluorescence signal from the vesicle indicates that the lipid membrane remains intact. Conversely, a signal decay (after controlling for photo-bleaching) suggests that the encapsulated fluorophore leaks out due to a compromised membrane. Monitoring peptide-induced leakage kinetics can reveal the “mode of action” of the peptide, which in turn can help interpret a particular permeabilization “mechanism”. In this vein, most previous studies used a suspension of large unilamellar vesicles (LUVs; diameters spanning 100–1000 nm) for monitoring peptide activity. In such cases, the user measures the average fluorescence values of the LUVs as a single population and hence such experiments provide information on the average leakage kinetics of the vesicle population as a whole. Consequently, the coarse-grained resolution of the extracted information only enables minimal interpretation of the mode of action—it does not allow the investigation of heterogeneity in the mode of action at the single-vesicle level. To enhance the quality and level of detail extracted from these experiments, giant unilamellar vesicles (GUVs), which can be studied individually, are used instead. In particular, negatively charged GUVs have been shown as excellent models for studying the efficacy of cationic peptides in Escherichia coli (E. coli) cells. We tailor the composition of PG lipids in our GUVs to approximately reflect the PG content in the inner membrane of E. coli. Note, here we are only considering the lipid headgroup composition to model the charge of the bacterial membrane. A recent study incorporating both a single-cell based approach (using E. coli) and the single-GUV based technology presented here confirmed that the membranolytic efficacy of the peptides correlated well with their corresponding antibacterial activity on E. coli, validating

the use of such membrane models for characterizing membrane active peptides. Typically, a handful of vesicles (diameters >1000 nm) can be directly visualized using fluorescence microscopy enabling the monitoring of the peptide-induced leakage kinetics of individual GUVs with a defined spatiotemporal resolution. However, such methods require laborious manual handling and suffer from a lack of control over experimental conditions and low throughput. To overcome these challenges, we previously devised a bespoke microfluidic platform that integrated high-throughput GUV formation and immobilization on-chip in parallel trapping arrays. GUV trapping and manipulation in microfluidic devices offers a powerful analytical tool for studying these membrane models in a highly controlled manner; a range of options are now available for such studies. While other microfluidic methods of GUV manipulation can achieve a similar level of control and throughput, such as continuous flow devices, these typically can study kinetic processes on the order of seconds to minutes, while peptide activity at relevant doses typically occurs on the order of hours. Our format enables the continuous administration of different peptides at a set dose in a parallelized manner on thousands of trapped vesicles. This microfluidic platform is thus perfectly suited to investigating the mode of action of membrane-active agents on a population of thousands of model membranes, while maintaining single-GUV and high spatiotemporal resolution.

In this work, we leverage our ability to monitor the response of thousands of individual GUVs to a range of AMPs; in terms of single GUVs analyzed per experiment, this is orders of magnitude greater than traditional techniques. We optimize a bespoke microfluidic platform into an the effective high-throughput device, dubbed a GUV studio, and demonstrate its applicability to elucidate antimicrobial modes of action. We relate the modes of action to the timing of dye leakage from the GUVs after applying a defined peptide dosage and with molecular mechanisms elucidated by AFM. By identifying and quantifying modes of disruption or leakage and correlating them with molecular mechanisms, we provide information that can inform appropriate dosage and administration regimens for antimicrobial agents. Our ability to track thousands of single GUVs in parallel thus delivers a novel analytical methodology for the understanding of the activity of membrane active peptides.

### EXPERIMENTAL SECTION

**Optofluidic Leakage Assay.** The optofluidic GUV leakage assay builds on our previous work studying the activity of cecropin B. The assay utilizes the GUV studio to standardize conditions and overcome numerous technical drawbacks that classical leakage techniques suffer from. The assay is designed to monitor the leakage kinetics of the encapsulated fluorophore pyranine from a population of GUVs at single-vesicle resolution, under continuous peptide administration. Our microfluidics platform facilitates both control over the dosed peptide concentrations and the imaging of hundreds to thousands of GUVs at desirable spatiotemporal resolution. Importantly, the continuous perfusion of the peptides ensures that all the GUVs studied are exposed to similar amounts of the peptides simultaneously, negating any potential inoculum effect. Another benefit of using continuous perfusion is that it minimizes the technical variations caused by non-specific binding of peptides to the walls of the device.
**GUV Studio.** As described in Figure 1, the GUV studio incorporates an on-chip GUV formation component that relies on the octanol-assisted liposome assembly (OLA) technique. The schematic outlines the platform’s capabilities in preparing GUVs, vesicle trapping, and controlled peptide administration. The GUV formation element is based on the OLA technique. The platform integrates eight separate chambers each encompassing an array of 720 vesicle traps. Vesicles are transported from the formation junction to the trap arrays via a second layer of microfluidic channels, labeled as the connector chip (in yellow). This design features a pneumatic valve system in a third layer underneath the main chip (in red). The valve system consists of three control channels that can be used to facilitate controlled switching between the different stages of the experiment. Insets show microscopic fluorescence images of vesicle formation (left) and trapping (right). In the lefthand inset, the membrane fluorescence is shown above and the fluorescence of the dye in the lumen is shown below. The lipid phase is tagged with Liss Rhod PE, and the GUVs encapsulate the dye HPTS. Scale bars represent 50 μm.

**GUV Preparation.** The inner aqueous (IA) solution of GUVs consisted of a PBS buffer (10 mM phosphate buffer, 2.7 mM KCl, and 137 mM NaCl) at pH 7.4, in addition to 200 mM sucrose and 15% v/v glycerol. The outer aqueous solution additionally contained 50 mg mL\(^{-1}\) of Kolliphor P-188 (Sigma-Aldrich, U.K.) and a metal connector tip (Gauge 23 blunt end, Intertronics). A neMESYS syringe pump with a 250 μL Duran borosilicate glass syringe (ILS, Germany) was connected to outlet A via an Upchurch 1520 G tubing (0.03” × 0.06”) (Figure 1). A more detailed description of the device operation and data acquisition can be found in the Supporting Information (Figure S1).

**Peptide Preparation.** NI01 and all its derivatives were synthesized, identified, and purified as previously published. Briefly, they were assembled in a Liberty microwave peptide synthesizer (CEM Corp.). All peptides were then purified by
and α the topography of NI01-treated PC/PG (3:1) SLBs mimicking bacterial membranes. The floral-shaped topography consists of bilayer spanning Crystal structure of NI01. Ribbon representation from the N terminus (green) to the C terminus (orange). (C) In-liquid AFM images representing NI01 primary structure, its 2-helix substructure derivatives, and the arginine mutant R-NI01. The three substructures are notated as (α1α2, α2α3, and α3α4). The four helices are color coded above the sequences and labeled α1−α4 accordingly. Turns are underlined in the sequences. (B) Crystal structure of NI01. Ribbon representation from the N terminus (green) to the C terminus (orange). (C) In-liquid AFM images representing the topography of NI01-treated PC/PG (3:1) SLBs mimicking bacterial membranes. The floral-shaped topography consists of bilayer spanning petal-shaped pores (dark regions) and a membrane thinning patch in the center (light regions).

Figure 2. Overview of the NI01 peptide series and the observed dual mechanism against anionic lipid bilayers.34 (A) Sequences of the epidermicin NI01 primary structure, its 2-helix substructure series, and the arginine mutant R-NI01. The three substructures are notated as (α1α2, α2α3, and α3α4). The four helices are color coded above the sequences and labeled α1−α4 accordingly. Turns are underlined in the sequences. (B) Crystal structure of NI01. Ribbon representation from the N terminus (green) to the C terminus (orange). (C) In-liquid AFM images representing the topography of NI01-treated PC/PG (3:1) SLBs mimicking bacterial membranes. The floral-shaped topography consists of bilayer spanning petal-shaped pores (dark regions) and a membrane thinning patch in the center (light regions).

The topographic imaging of SLBs in aqueous buffers was performed on a Multimode 8 AFM system (Bruker AXS, USA) using Peak Force Tapping mode and MSNL-E cantilevers (Bruker AFM probes, USA). Images were taken at a Peak Force frequency of 2 kHz, an amplitude of 10−20 nm, and a set point of 10−30 mV (<100 pN). NI01 or its derivatives were introduced into a 10 μL fluid cell (Bruker AXS, USA) to the intended final concentration. We note that the lipids used for the AFM and GUV studies have similar headgroups but differ in the structure of the hydrocarbon tail. Although it is generally considered that the lipid composition may impact on membrane disruption, the differences in headgroups are the critical determinants, and these are PC/PG mixtures for both studies. Lipids with mixed, unsaturated, and saturated aliphatic chains produce similar results when it comes to membrane disruption in GUVs and live bacteria alike.35 Unlike pore-forming proteins, membrane-active antibiotics are not limited to the formation of conserved pore structures that occur in particular membranes. Instead, they porate different bacteria (including Gram-negatives and Gram-positives) whose membrane compositions vary. In the context of this work, we screen the kinetics of membrane disruption in a statistically significant manner (using GUVs) and correlate this with the patterns observed by AFM, which have previously been correlated with antimicrobial activities.34

■ RESULTS AND DISCUSSION

We employed an optofluidic leakage assay to monitor single-vesicle leakage events (LEs) utilizing the GUV studio depicted in Figure 1. The platform allows the user to continuously dose several populations of GUVs (each containing a membrane impermeable dye) with an antimicrobial agent at different concentrations in parallel over 7.5 h. Furthermore, due to the distinct handling advantages of the microfluidic GUV studio platform,23 all GUV populations within the experiment possessed excellent homogeneity of size and composition. Indeed, in a direct comparison between OLA generated and
electroformed vesicles, we previously showed how OLA results in a more homogeneous vesicle population than electroformation. A vesicle LE is defined as the >50% loss of fluorescent dye from the vesicle lumen. Under our definition of LEs, we do not discriminate between rapid vesicle rupture and slow leakage, although our device is certainly able to discriminate between rapid rupture and slow leakage on the order of several minutes. Furthermore, we note that leakage of the dye may be due to detergent-like rupture of the GUV membrane, or other mechanisms such as pore formation. The width (in time) of the transition from high to low vesicle fluorescence intensity in the heat maps (i.e., the portion of the heat map in white) informs on the time of leakage of the dye from the vesicle lumen. Importantly, within each experiment (i.e., a single run of the GUV studio device), all the trapped populations of GUVs were assembled on the same device. The source of all the GUVs in an experiment was hence identical. To allow direct comparisons, the experimental parameters and conditions were identical for all the peptides studied, with peptide sequence being the only variable.

To demonstrate the approach, two series of representative AMPs were selected. One series consisted of bacteriocin-derived peptides including a four-helix bacteriocin epidermicin (NI01), its all-D analogue (D-NI01), an arginine mutant (R-NI01), and three 2-helix substructure derivatives (Figure 2). Another series was comprised of well-studied host defense peptides discovered in different organisms, including melittin (a bee venom peptide), an amphibian peptide magainin 2 (from Xenopus laevis), an antibiotic alamethicin from the fungus Trichoderma viride, a last-resort antibiotic polymyxin B (from Bacillus polymyxa), and a human β-defensin (hβD-3-1). The response of a GUV population to defined concentrations of each peptide was monitored with single GUV resolution. Four different peptide concentrations were run in parallel to probe concentration dependent effects. The mode of action was determined by measuring the time distribution of single GUV LEs and the percentage survival of a GUV population at the end of an experiment as summarized in the corresponding heat maps.

**NI01 Mode of Action.** NI01 exhibits a synergistic multiple mechanism in anionic phospholipid bilayers characterized by the formation of transmembrane lesions and pores and membrane thinning patches, ~2 nm in depth half-way through the bilayer, in a floral pattern (Figure 2C). The mechanism of NI01 has also been shown to be non-stereoselective as its all-L and all-D forms demonstrated similar membrane disruption patterns. Given this mechanism, we sought to examine the modes of action for both the all-L and all-D forms of NI01 (Figure 3). First, we found a strong dependence of GUV survival on the peptide concentration when treated with 5 μM of the peptide, the vast majority (~89.6%) of the GUVs survived treatment (i.e., no dye leakage was observed) over the timescales measured. However, for higher concentrations (10 μM and above), the entire GUV population showed complete leakage of the encapsulated dye molecules within the timescales of the experiment, with the mean time of leakage decreasing with increasing concentration. The large fraction of vesicle survival when dosed with 5 μM of the peptide may result from the peptide not reaching the threshold concentration necessary to induce leakage in individual vesicles. For the remaining peptide dosages, LEs converged into a unimodal narrow distribution around a concentration-specific timepoint (LE breakdown is provided in the Supporting Information). The LEs’ mean time point inversely correlated with the corresponding peptide concentration (10 μM: 322.2 ± 44.0 (std. dev.) min N = 388, 25 μM: 136.2 ± 34.6 min N = 469, 50 μM: 74.4 ± 14.8 min N = 599) (Figure S3). This indicates that NI01 is not a stochastic membrane disruptor in GUVs because its action is concentration dependent, and the LEs are monodisperse. The heat maps for the all-L form experiments demonstrate an almost identical response to the all-L form (Figure 3B). Therefore, both the all-L and all-D forms exhibit similar modes of action consistent with their mechanisms established by AFM.
Dissecting NI01 to Understand the Relation between Its Mechanism and Mode of Action. The observed dual mechanism of the four-helix bacteriocin was further examined by deconstructing the peptide into three 2-helix hairpin substructures and monitoring the modes of action for each (Figure 4A). Each of the hairpins supports a specific part of NI01’s multiple mechanisms: the $\alpha_1\alpha_2$ and $\alpha_2\alpha_3$ hairpins form elongated transmembrane lesions and membrane thinning patches, respectively (Figure S7). In contrast, $\alpha_3\alpha_4$ combines the formation of thinning patches with that of circular transmembrane pores. We therefore probed whether the differences in mechanisms for the three hairpins translate to different modes of action. The experiments were conducted at four different concentrations for each substructure. LEs were observed at the 50 $\mu$M concentration (Figure 4). At the lower doses, most of the GUVs survived the treatment within the timescales measured. Compared to the data for NI01 (Figure 3), this indicates that the efficacy of the substructures against GUVs is lower than the activity of the full-size NI01. At the 50 $\mu$M concentration for the $\alpha_1\alpha_2$ and $\alpha_3\alpha_4$ hairpins, we observed the total leakage of all the individual vesicles. The timing of the LEs grouped into a unimodal narrow distribution around a similar timepoint ($240.1 \pm 10.7$ min $N = 477$ and $267.4 \pm 10.8$ min $N = 408$) for both peptides, respectively. With regard to $\alpha_2\alpha_3$, the recorded timing of LEs showed some stochasticity and did not converge around a single time point. At the end of the experiment (450 min), the population was split into $\sim45\%$ survivors and $\sim55\%$ disrupted vesicles (Figure S4). These results highlight a correlation between the membrane active mechanism of a certain peptide with its mode of action as defined via a signature LE time distribution. Peptides that introduce transmembrane lesions have similar modes of action on populations of GUVs. With regards to the mechanisms of NI01 and $\alpha_3\alpha_4$, the mode of action appeared to be dictated by transmembrane poration as the dominant mechanism over membrane thinning. Similar to NI01, the substructures do not qualify as stochastic membrane disruptors in GUVs because peptide action is concentration dependent, and LEs only occur after a certain lag time.

Exploring Correlations between the Mechanism and Mode of Action of NI01. Electrostatic interactions are one of the main driving forces responsible for the membrane activity of AMPs. To gain a better insight into the role of charge in the activity of NI01, an all-arginine mutant of NI01 was synthesized, in which all cationic residues in the native sequence were replaced with arginines to ensure stronger electrostatic interactions with anionic membrane surfaces. This difference in membrane interaction is believed to arise from the fact that arginine is positively charged at all stages of membrane insertion, whereas lysine can become deprotonated once in the membrane. Note that in addition to the charge differences, structural changes may have an impact on the disruption pattern because the arginine sidechain has a higher molecular weight and volume than the lysine sidechain. Similar to the other peptide, the mode of action for the mutant was probed using four different peptide concentrations (5, 10, 25, and 50 $\mu$M). A high percentage of LEs was observed even at the 5 $\mu$M concentration indicating that the efficacy of the mutant against GUVs appeared higher than that of the original NI01 structure (survival: NI01 = 89.6%, R-NI01 = 0%).

Figure 4. Dissecting multiple mechanisms of NI01 into substructure derivatives and corresponding modes of action. (A) Three 2-helix substructures derived from NI01. (B) Schematics representing the inner and outer leaflets of a single lipid bilayer and the corresponding mechanisms of membrane thinning or transmembrane lesions for each of the substructures. The schematic is based on the interpreted data from the in-liquid AFM topography images of anionic PC/PG (3:1) SLBs treated with the two-helix hairpins (AFM images shown in Figure S7). (C) Heat plots mapping the modes of action of the three substructures, $\alpha_1\alpha_2$, $\alpha_2\alpha_3$, and $\alpha_3\alpha_4$ against anionic PC/PG (3:1) GUVs.
Furthermore, the distribution of timepoints of vesicle LEs at the lower peptide concentrations proved to be wide (5 μM: 122.6 ± 90.4 min N = 440, 10 μM: 102.8 ± 76.2 min N = 339, and 25 μM: 134 ± 74.4 min N = 479). Increasing the peptide concentration led to a second peak and the convergence of LEs clustering around a specific, concentration-dependent time point (50 μM: 103.0 ± 32.3 min N = 518) (Figure S5). This suggests that at 50 μM, a threshold concentration is reached. At this concentration, LEs clustered into a unimodal narrow distribution at the mentioned timepoint. For R-NI01, thinning patches were the mechanism identified via AFM measurements, with no detection of the transmembrane pores that were observed with the original NI01 peptide (Figure S7).

Given this, it is interesting that, as mentioned above, the vesicle studies revealed strong membrane compromising activity even with 5 μM of the mutant peptide, in comparison to the original NI01 peptide at this concentration. The fact that R-NI01 has a strong effect on the GUVs is, however, consistent with the stronger peptide-lipid interactions furnished by arginine residues. Unlike lysine, arginine provides five hydrogen-bond phosphate clusters, which enhance the affinity of the peptide to phospholipids changing the dynamics of membrane disruption to favor membrane exfoliation. However, comparing the time distributions of LEs between NI01 and R-NI01 (Figure S5B), we can say unequivocally that, as far as the mode of action is concerned, R-NI01 qualifies as a stochastic membrane disruptor in GUVs because LEs occur without any specific lag period and are spread over a wide range of time points (Figure S5). The heterogeneity of GUV responses to R-NI01 is further displayed in Figure S2.

**Exploring Model Mechanisms and Correlating Modes of Action.** To further explore correlations between mechanisms of membrane disruption and modes of action, we investigated a series of archetypal and well-studied AMPs. These were chosen to allow for a comparison to the inferences drawn from the NI01 series. These peptides have different molecular mechanisms and modes of action. Melittin and magainin 2 are known to form toroidal pores with the distinction of having graded and all-or-none leakage characteristics, respectively. Graded action refers to the mode, where all the individual vesicles show similar leakage characteristics and hence the time points of LEs are tightly distributed (unimodal). On the other hand, all-or-none action indicates that the population can be split into intact and leaked vesicles (bimodal). Alamethicin is postulated as a model peptide for barrel-stave pore formation. Barrel-stave and toroidal pores are both transmembrane. However, the walls of barrel-stave pores only consist of peptides, while toroidal pores cause the membrane to bend, with the phospholipid headgroups facing the interior of the pore. Polymyxin B lyses membranes in a detergent-like manner, while the mechanism of the linear form of hBD-3 is believed to involve disordered toroidal pore formation. The concentration at which these peptides were studied, the lipid composition of reconstituted membranes used in the studies, and other experimental parameters are factors that may affect their activity characteristics. With this in mind, we characterized the modes of action of these peptides using the GUV studio under the conditions used for the first series, to probe unique signatures for the LE time distributions for these peptides (Figures 6 and S6). We then correlate the LE time distribution with the modes of action of the peptide against their molecular mechanisms.

First, we compared the LE time distributions of melittin and magainin 2. For melittin, we detected the total leakage of the entire vesicle population at all concentrations but 5 μM—a scenario reminiscent of NI01’s mode of action. For magainin 2, LEs corresponding to most of the vesicle population were observed only at 25 and 50 μM. Moreover, at 10 μM, the peptide displayed a unique LE time distribution, where a ~44% subpopulation of vesicles remained intact throughout peptide treatment. With the process of vesicle leakage being stochastic, the time distribution of LEs that took place converged around a certain time point (10 μM: 116.1 ± 77 min N = 177) suggesting that magainin 2 forms transmembrane pores, which is in contrast to the effects observed for α2α3 characterized by the mechanism of membrane thinning. These results concur with those reported by others for melittin and magainin 2. Although both peptides by virtue of forming transmembrane pores exhibit a similar molecular mechanism, our results suggest that their modes of action are different in that melittin shows a graded characteristic, while magainin 2 favors the all-or-none characteristic. We then compared melittin with alamethicin. Both peptides gave comparable LE time distributions at different concentrations, similar to what was observed for NI01. The majority of LEs occur after a certain lag time (10 μM: 165.3 ± 62.6 min N = 406, 25 μM: 61 ± 17.9 min N = 641, and 50 μM: 34.1 ± 23.8 min N = 645), which inversely correlated with corresponding peptide concentrations, thus indicating that both the toroidal and barrel-stave poration follow the two-state model. Next, our assay revealed that polymyxin B was the most potent among all of the tested peptides. The total leakage of all the vesicles in the population was observed at all
The LEs do not occur after a specific lag period, the latter time for the dye to leak out of an individual vesicle. Because increasing the duration of vesicle leakage (i.e., the amount of 153 min $\pm 569$, 10 $\mu$M: 388, 25 $\mu$M: 28.6 $\pm$ 35.9 min $N = 480$, and 50 $\mu$M: 11.5 $\pm$ 16.9 min $N = 430$) supporting the previous two-state model, as with increasing peptide concentration the lag time duration allowed by the GUV studio offer an effective approach to systematically characterize membrane-active antimicrobials. Assigning a mode of action, which reflects the kinetics of peptide-membrane interactions and may provide correlations with the underlying molecular mechanisms, is less common. We designed and applied a bespoke GUV studio to determine the modes of action of a range of AMPs. The study provided important insights into correlations between the molecular mechanisms of the peptides and their modes of action. First, for the all-L and all-D stereoisomeric forms of a peptide demonstrating the same molecular mechanism, the mode of action is also the same. Second, if a peptide exhibits multiple mechanisms, the main determinant of its mode of action is its most disruptive mechanism. Third, LEs caused by transmembrane poration follow one of two modes of action. In one, a two-state mode, pore formation ensues upon reaching a threshold peptide concentration. This mode is characterized by LEs occurring within a narrow time range, with peptide concentration determining the mean time point of the LEs. Another one is a spontaneous mode, where LEs can take place before reaching a threshold peptide concentration. This leads to LEs that are distributed over a wide range of time points, for which the impact of the peptide concentration is limited to the width of the LE time distribution. Fourth, pore-forming peptides demonstrating different molecular mechanisms, for example, melittin and alamethicin, can share a similar mode of action. By contrast, peptides exhibiting the same molecular mechanism, for example, melittin and magainin 2, can demonstrate different, for example, “graded” or “all-or-none”, modes of action. Finally, GUV responses to peptides inducing membrane thinning typically correlate with a mode of action that is associated with stochastic LEs. The majority of the vesicle population in this scenario remains intact. The demonstrated correlations between the molecular mechanisms of peptides and their kinetic modes of action suggest that the sole use of MIC measurements to define antimicrobial treatments requires reconsideration. The direct comparisons allowed by the GUV studio offer an effective approach to systematically characterize membrane-active antimicrobials. We also note that we can in principle change the GUV membrane composition to instead more closely resemble mammalian cell membranes and thereby use the platform to investigate the potential cytotoxicity of these compounds. Thus, the platform could inform on both the antimicrobial efficacy and the safety profile of candidate AMPs. By expanding this methodology to a larger number of peptides representing different antimicrobial mechanisms, it will be possible to systematically characterize their different modes of action in a standardized manner, which may better inform the design and translation of clinically acceptable peptides.

**ASSOCIATED CONTENT**

*Supporting Information*

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.analchem.1c03564.

Device fabrication and operation of the GUV studio and the protocol for data analysis; visualization of leakage heterogeneity in vesicle populations; histograms of
leakage event distributions for all peptides; summary of leakage event distributions including mean LE time-point, standard deviation, and survival; and AFM images (PDF)

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Notes
The authors declare no competing financial interest.

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