Mechanisms of bacterial membrane permeabilization by crotalicidin (Ctn) and its fragment Ctn(15–34), antimicrobial peptides from rattlesnake venom

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Crotalicidin (Ctn), a cathelicidin-related peptide from the venom of a South American rattlesnake, possesses potent antimicrobial, antitumor, and antifungal properties. Previously, we have shown that its C-terminal fragment, Ctn(15–34), retains the antimicrobial and antitumor activities but is less toxic to healthy cells and has improved serum stability. Here, we investigated the mechanisms of action of Ctn and Ctn(15–34) against Gram-negative bacteria. Both peptides were bactericidal, killing ~90% of Escherichia coli and Pseudomonas aeruginosa cells within 90–120 and 5–30 min, respectively. Studies of ξ potential at the bacterial cell membrane suggested that both peptides accumulate at and neutralize negative charges on the bacterial surface. Flow cytometry experiments confirmed that both peptides permeabilize the bacterial cell membrane but suggested slightly different mechanisms of action. Ctn(15–34) permeabilized the membrane immediately upon addition to the cells, whereas Ctn had a lag phase before inducing membrane damage and exhibited more complex cell-killing activity, probably because of two different modes of membrane permeabilization. Using surface plasmon resonance and leakage assays with model vesicles, we confirmed that Ctn(15–34) binds to and disrupts lipid membranes and also observed that Ctn(15–34) has a preference for vesicles that mimic bacterial or tumor cell membranes. Atomic force microscopy visualized the effect of these peptides on bacterial cells, and confocal microscopy confirmed their localization on the bacterial surface. Our studies shed light onto the antimicrobial mechanisms of Ctn and Ctn(15–34), suggesting Ctn(15–34) as a promising lead for development as an antibacterial/antitumor agent.

New antimicrobial drugs are urgently needed to address the growing challenge of bacterial resistance to existing antibiotics. Misuse of classical antibiotics has increased the number of superbugs and created a critical situation whereby previously controlled pathogens could in the future cause major morbidity or mortality (1, 2). This alarming growth of multidrug-resistant pathogens has prompted an intensive search for anti-infective drugs with novel mechanisms of action (3, 4). In particular, antimicrobial peptides (AMPs) have emerged as promising alternatives due to their broad-spectrum activity (including superbugs), selectivity, and mechanisms of action that potentially hinder the development of resistance (5).

AMPs are ancient weapons of the host defense machinery, present in all life domains (6). Although they can act in several possible ways to accomplish microbial cell death (e.g. membrane disruption, apoptosis induction, and internal target inhibi...
bition) (7, 8), an initial common step in the process is their recruitment onto the bacterial cell surface (9, 10). Accordingly, most AMPs display fairly conserved structural and physicochemical properties, such as positive net charge, high content of hydrophobic amino acid residues, or amphipathic structure, all favoring interaction with and insertion into membranes (11, 12). Cathelicidins are a large family of AMPs whose unifying feature is the presence of a conserved cathelin (cathepsin L inhibitor) domain at the N terminus. In contrast, their C-terminal domains contain a mature and active AMP and display high inter- and intraspecies diversity (13). Cathelicidins have been shown to be active against a broad range of targets, including bacteria, enveloped viruses, and fungi (14). In addition to causing direct pathogen killing, cathelicidins can modulate the immune response by assisting with pathogen clearance (15). Cathelicidins have been isolated from a wide range of organisms, including mammals (16–18), birds (19), fish (20), frogs (21), and marine (22) and terrestrial snakes (23–25).

We recently identified a new family of cathelicidin-like peptides named vipericidins in the venom glands of various South American pit viper snakes and experimentally validated them as AMPs (26). Crotalicidin (Ctn), the most active vipericidin, is a 34-residue helical peptide found in Crotalus durissus terrificus and was selected for further study. Ctn displays powerful antimicrobial, as well as antitumor and antifungal, activity. However, it is moderately hemolytic and unstable in serum (27, 28). To overcome these limitations, a rational dissection of the Ctn sequence was undertaken, aimed at identifying active motifs with enhanced properties. The search produced as main lead Ctn(15–34), a structurally disordered 20-mer that spans the C terminus and replicates the antibacterial and antitumor activities of the parental peptide but is less toxic toward healthy cells and significantly more stable in human serum (27). Investigating the mechanism of action on bacteria of both Ctn and Ctn(15–34) is of considerable interest, particularly given the potential therapeutic applications of the latter peptide.

In this study, the mode of action of Ctn and Ctn(15–34) toward two Gram-negative bacterial species was investigated in detail. We determined bactericidal concentrations and quantitative kinetics of cell death and measured peptide accumulation on bacterial cell surfaces. Combining flow cytometry and colony count procedures, we established that bacterial death is accomplished by membrane disruption. We also monitored peptide uptake and membrane permeabilization in real time to ascertain whether both effects are achieved simultaneously or peptide pre-accumulation precedes loss of viability. To visualize peptide localization and structural damage on bacteria surface, we carried out confocal and atomic force microscopy experiments. Furthermore, to gain insights into the cell selectivity and the ability to disrupt lipid bilayers of the fragment Ctn(15–34), we performed experiments with model vesicles mimicking healthy human and bacterial cell membranes. Altogether, our results provide strong evidence that Ctn and Ctn(15–34) act by inducing disruption of the bacterial cell; they also exemplify how a set of optimized methodologies can be combined to evaluate the action of AMPs at the membrane level.

### Results

#### Synthetic peptides

The amino acid sequences in Table 1 were prepared in C-terminal amide form by Fmoc (N-(9-fluorenyl)methoxycarbonyl) solid-phase synthesis and purified to >95% purity (see Fig. S1 for HPLC and MS data). For the N-terminal rhodamine B (RhB)-labeled peptides, the two peaks observed in the chromatograms of purified compounds (Fig. S1a) are due to RhB atropisomerism. The overall hydrophobicity of unlabeled and RhB-labeled peptides can be compared by the percentage of acetonitrile (ACN) at which they elute in HPLC (Table 1). Values for unlabeled Ctn and Ctn(15–34) were 30.9 and 30.3% ACN, respectively, whereas RhB-labeled versions eluted at 34.7 and 34.2% ACN, respectively, underlining the increase in hydrophobicity brought about by RhB labeling.

#### Antibacterial and bactericidal activity

We evaluated the antimicrobial activity of Ctn and Ctn(15–34) against two Gram-negative bacterial strains: Escherichia coli ATCC 25922 (E. coli) and Pseudomonas aeruginosa ATCC 27853 (P. aeruginosa). As shown in Table 2, Ctn has the lowest minimal inhibitory concentration (MIC) for E. coli as well as for P. aeruginosa (0.78 and 1.56 μM, respectively). Ctn(15–34) also has a low MIC for E. coli (3.13 μM), but the concentration required to prevent P. aeruginosa visible growth is higher (12.5 μM). To determine whether the peptides were bactericidal or bacteriostatic, the minimal bactericidal concentrations (MBCs) were also determined (Table 2). The data show both Ctn and Ctn(15–34) to be bactericidal, as MBCs exceed MICs by <2-fold (29). The MIC/MBC determinations reveal that more peptide is required for bacterial death when higher bacterial inocula are used. This difference is significant for E. coli, for

### Table 1

Peptides used in this study

| Peptide | Amino acid sequence | Theoretical mass | Experimental mass | Purity | ACN gradient | Retention time | ACN |
|---------|---------------------|------------------|------------------|-------|--------------|---------------|-----|
| Ctn/    | KRFKKFKVKSVKKRIKFEKPMVIGTVIPF-amide | 4151.39 | 4152.45 | 98 | 15–40 | 9.556 | 30.9 |
| RhB-Ctn | RhB-KRFKKFKVKSVKKRIKFEKPMVIGTVIPF-amide | 4576.94 | 4575.97 | 99 | 20–40 | 10.997 | 34.7 |
| Ctn(15–34) | KKKRLKKIFKKPMVIGTVIPF-amide | 2371.11 | 2371.23 | 99 | 10–40 | 10.131 | 30.3 |
| RhB-Ctn(15–34) | RhB-KKKRLKKIFKKPMVIGTVIPF-amide | 2796.66 | 2795.90 | 98 | 20–40 | 10.670 | 34.2 |

* Theoretical molecular mass was calculated using GPMAW version 8.10.

* Experimental molecular mass was determined from the MS spectra shown in Fig. S1b.

* Peptide purity was estimated by peak integration of the analytical HPLC chromatograms shown in Fig. S1a.

* ACN gradient over 15 min used to run the analytical HPLC. Range indicates the initial and final ACN concentrations.

* Percentage of ACN corresponding to the eluted peptide, calculated from the gradient used and the retention time.

* Protein Data Bank accession code of Ctn: 2MW7.
Membrane-disruptive effect of Ctn and Ctn(15–34)

Table 2
MIC and MBC of Ctn and Ctn(15–34) against two standard Gram-negative bacterial strains

| Peptide      | E. coli ATCC 25922 | P. aeruginosa ATCC 27853 |
|--------------|-------------------|-------------------------|
|              | 5 × 10^5 cfu/ml   | 10^5 cfu/ml             |
|              | 5 × 10^5 cfu/ml   | 10^5 cfu/ml             |
| MIC (µM)     |                   |                         |
| Ctn          | 0.78              | 6.25                    |
| Ctn(15–34)   | 3.13              | 50                      |
| MBC (µM)     |                   |                         |
| Ctn          | 1.56              | 6.25                    |
| Ctn(15–34)   | 6.25              | 50                      |

which up to 3 times more peptide is needed. These results support previous observations suggesting that the therapeutic effect of AMPs depends on the peptide/cell ratio rather than directly on peptide concentration (30, 31). In tune with this suggestion, and to avoid the results being influenced by the use of different peptide/cell ratios, peptide concentrations for mechanism-of-action studies were adjusted to the bacterial inoculum used for each specific experiment.

Time-dependent bacterial death

The killing kinetics of E. coli and P. aeruginosa by Ctn and Ctn(15–34) at their respective MBCs was evaluated using a time-kill assay. The results in Fig. 1a show that for both peptides, a bactericidal effect (99.9% reduction of bacterial viability) against E. coli is observed within 4 h. At 90 min, Ctn kills >90% of bacteria, whereas Ctn(15–34) needs 120 min for a similar effect. Ctn and Ctn(15–34) are faster at killing P. aeruginosa, requiring only 5 or 30 min for 90% cell death, respectively, with bactericidal effect observed at 15 and 90 min, respectively. This relatively fast effect against E. coli, even faster against P. aeruginosa, suggests that both peptides act by direct damage to the bacterial membrane rather than by inhibition of an internal target.

Membrane surface charge neutralization

To investigate whether surface charge can be neutralized by the accumulation of the peptides at the membrane of E. coli or P. aeruginosa, ζ potential measurements were carried out. The results show that, as expected (32, 33), in the absence of peptide, both bacteria possess a negative potential (Fig. 1b, ~−24 mV for E. coli and ~−21 mV for P. aeruginosa). In the presence of increasing peptide concentrations, a shift to positive ζ potential values is observed. In E. coli, complete neutralization of the surface charge occurs at 0.78−3.13 µM Ctn and 1.56−3.13 µM Ctn(15–34), whereas for P. aeruginosa, total surface charge neutralization is reached at 0.78−3.13 µM Ctn and 3.13−6.25 µM Ctn(15–34). For both bacteria, membrane neutralization occurs at concentrations below the MIC/MBC of either peptide.

Membrane permeabilization and viability correlation studies

To clarify whether E. coli and P. aeruginosa death induced by Ctn and Ctn(15–34) is due to direct membrane disruption, correlation between membrane permeabilization and bacterial viability was studied. To this end, a flow cytometry assay (FCA) with SYTOX® Green to examine membrane permeabilization and a plate colony count to quantify viable cells were performed. As a cell-impermeable nucleic acid stain that only labels bacteria with compromised plasma membrane, SYTOX® Green is a useful tool in bacterial viability studies. As shown in Fig. 2, an increased percentage of bacterial cells with SYTOX® Green-permeable membrane is accompanied by a decrease in cell viability by the colony count method. FCA histograms on the SYTOX® Green channel (Fig. 2a) shift to higher fluorescence intensities indicative of E. coli and P. aeruginosa membrane disruption with increasing peptide concentration. Negative and positive controls (live and dead bacteria; black and gray histograms, respectively) were used to define gates and calculate permeabilization percentages, shown in Fig. 2b along with the viability percentages obtained in the colony count assay. A direct correlation between both sets of data (Fig. 2b) confirms that antibacterial activity of Ctn and Ctn(15–34) is mediated by membrane permeabilization.

To correlate the membrane permeabilization effect with peptide uptake (including both membrane-bound and intracellularly internalized), a parallel FCA experiment was performed with RhB-labeled versions of both peptides. From the SYTOX® Green and RhB channel readings (Fig. S2, a and b), the percentages of permeabilized and of peptide-internalizing cells, respectively, were determined (Fig. S2c). For both Ctn and Ctn(15–34), an increase in E. coli permeabilization coincides with increased peptide uptake, suggesting a direct correlation between the membrane damage observed and peptide entry/binding. The dose–response curve for RhB-Ctn (Fig. S2c, continuous red line) is similar to that for unlabeled Ctn (Fig. 2b, continuous red line), suggesting that RhB does not interfere with the peptide effect. In contrast, the profiles for RhB-labeled (Fig. S2c, continuous blue line) and unlabeled Ctn(15–34) (Fig. 2b, continuous blue line) are different. Whereas the latter induces 50% permeabilization in the 6.25–12.5 µM range, the percentage rises to 80% for the RhB-labeled analogue at similar concentrations. Moreover, at higher (25–100 µM) concentrations, a decrease in membrane permeabilization and uptake of RhB-Ctn(15–34) occurs, an effect not observed for unlabeled Ctn(15–34). This probably reflects a loss in integrity of the bacterial cell, no longer capable of retaining the labeled peptide within, and with its DNA (SYTOX® Green target) either damaged or leaking out. The observation of N-terminal RhB label interfering with the activity of Ctn(15–34) calls for caution in extrapolating results from labeled to unlabeled versions of this peptide.
Membrane-disruptive effect of Ctn and Ctn(15–34)

Time-resolved peptide uptake and bacterial death

Changes in membrane permeabilization and peptide uptake were monitored by a time-resolved flow cytometry assay, which allows a more accurate study of the kinetics of peptide effect, compared with kinetic studies based on end-point sampling. We studied the membrane permeabilization of \( E. \ coli \) using SYTOX\(^\text{G}\) Green dye right after Ctn or Ctn(15–34) addition, monitoring the changes from negative to positive gates for 90 min. Fig. 3 (a and b) shows time-course results and negative and positive controls (for the whole acquisition, see Movies S1a and S1b for Ctn and Ctn(15–34), respectively). As detailed under “Experimental procedures,” data from the FCA histograms were used to generate the kinetic curves on Fig. 3c. Data were fitted using the two-state kinetic model described previously for other AMPs (34), assuming that peptide-bacteria interaction consists of an initial binding step, followed by permeabilization of the membrane.

Although both peptides induce permeabilization of \( \sim 90\% \) of bacteria after 90 min (Fig. 3c), the process differs between peptides at the early stages; based on the tendency of the experimental data, Ctn(15–34) seems to start killing bacteria right after its addition, and the permeabilization takes place in a single process. In contrast, Ctn shows an initial lag phase. Fitting of the kinetic data to a two-state model results in statistically poor residuals for some time intervals (Fig. S3), suggesting that for both peptides, permeabilization is more complex than assumed by the model. An additional slow event leading to permeabilization is suggested because the data deviate from the fit at longer times. In contrast, the fit is quite acceptable at shorter times. \( k_0 \), the membrane attachment rate constant, is higher for Ctn(15–34) than for Ctn (\( 2 \times 10^{-2} \) s\(^{-1}\) versus \( 1 \times 10^{-3} \) s\(^{-1}\)) as well as the permeabilization rate constant (\( 1 \times 10^{-3} \) s\(^{-1}\) versus \( 9 \times 10^{-4} \) s\(^{-1}\)). Cooperativity is not observed in either case (\( \beta \sim 0 \)), contrary to other AMPs (34).

To determine whether permeabilization is concomitant with internalization, kinetic studies with RhB-labeled peptides were run. Correlograms (Fig. S4a) at different time points show the changes recorded on the SYTOX\(^\text{G}\) Green and RhB channels (see Movies S2a and S2b for the entire acquisition of RhB-Ctn and RhB-Ctn(15–34), respectively). Kinetic curves (Fig. S4c) show both peptides becoming gradually bound to and/or internalized into the cells until an equilibrium is reached, but the uptake of RhB-Ctn(15–34) is faster than that of RhB-Ctn. Interestingly, RhB-Ctn(15–34) uptake precedes bacterial membrane damage, suggesting that peptide accumulation up to an effective threshold is necessary to trigger membrane permeabilization. In contrast, RhB-Ctn uptake shortly precedes membrane permeabilization, confirming that membrane becomes permeabilized as Ctn molecules are recruited to the surface.

Peptide distribution in bacteria

To investigate how peptides are localized within the bacteria, confocal microscopy was done on \( E. \ coli \) after incubation with RhB-labeled Ctn and Ctn(15–34) for 90 min. Images were also acquired using the SYTOX\(^\text{G}\) Green dye, to correlate peptide location with bacterial membrane permeabilization. Single-cell images acquired using the superresolution Airyscan mode are shown in Fig. 4a. Both peptides are preferentially localized around the bacteria, probably on the membrane. In some images, it was also possible to observe both peptides apparently...
co-localizing with SYTOX® Green (as shown in the fluorescence distribution plot; Fig. 4a). This suggests partial internalization and binding to DNA by the peptide, but the finding is not general for all bacteria. Images acquired in the normal confocal mode (Fig. S5) show untreated controls and bacteria treated with non-labeled peptides.

**Imaging peptide effect on bacterial cells**

To visualize structural changes on *E. coli* induced by Ctn and Ctn(15–34), individual bacteria were imaged by AFM after incubation with peptides at their MBC. As shown in Fig. 4b, both peptides induce major morphological alterations. Compared with control, the typical shape of the bacteria is lost, and substantial shrinkage, particularly at the central region, is observed. Additionally, treatment with either peptide also induces leakage of cytoplasmic content, as shown by images of bacterial surroundings. Cross-line profiles on Fig. 4b allow to visualize the location of three (for Ctn) or two (for Ctn(15–34)) protuberances in the septal poles of the bacteria and the aforementioned shrinkage at the middle.

**Ability of Ctn(15–34) to target the bacterial surface**

Gram-positive bacteria have lipoteichoic acid (LTA) at the membrane surface, whereas Gram-negative bacteria have lipopolysaccharide (LPS) (35). Both LTA and LPS are negatively charged and likely to be involved in the first contact of peptides with bacterial cells. To examine whether Ctn(15–34) and its labeled analogue RhB-Ctn(15–34) are able to bind to these molecules, we have used a *Limulus* amebocyte lysate (LAL) assay, which detects the presence of non-neutralized LTA and LPS (36). Results in Fig. 5a show that RhB-Ctn(15–34) neutralizes both LPS and LTA, whereas Ctn(15–34) is less efficient in doing it, especially for LTA.

**Interaction of Ctn(15–34) with model membranes**

The ability of Ctn(15–34) to bind model membranes composed of different lipid mixtures was examined by surface plas-
mon resonance (SPR). Previous studies suggested a preferential binding of Ctn(15–34) to bacteria and cancer cells rather than healthy eukaryotic cells (27). In view of this, a number of phospholipid compositions mimicking various types of cell membranes were assayed. Vesicles composed of 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC), a zwitterionic phospholipid forming neutral, fluid lipid bilayers at 25 °C, were compared with membranes made of POPC/sphingomyelin (SM)/cholesterol (Chol) (POPC/SM/Chol; 27:33:40 molar ratio), which forms bilayers with the liquid-ordered phase. POPC mimics the neutral fluid portions, whereas POPC/Chol/SM simulates the more-ordered raftlike portions of the outer leaflet in healthy human cells (37, 38). The inner leaflet of healthy human cell plasma membranes possesses zwitterionic phosphatidylethanolamine (PE)-, anionic phosphatidylserine (PS)-, and phosphatidylcholine (PC)-phospholipids (37, 38); thus, the mixture POPC/1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-L-serine (POPS)/1-palmitoyl-2-oleoyl-sn-glycero-phospho-L-ethanolamine (POPE) (60:20:20) has been included to mimic the inner leaflet of healthy cell membranes. Tumor cells possess larger proportions of negatively charged phospholipids exposed to the outer leaflet than healthy cells; therefore, the potential effect of negatively charged phospholipids was further examined by testing the affinity to membranes composed of POPC/POPS in 80:20 or 60:40 proportions. Finally, bacterial cell membranes have high negative charge and large proportions of phosphatidylglycerol (PG)-phospholipids; to mimic them, vesicles composed of POPC/POPG (80:20 or 60:40) or of an *E. coli* polar extract that contains a mixture of PE-phospholipids, PG-phospholipids, and cardiolipin were prepared.

**Figure 3. Time-resolved bacterial membrane permeabilization.** Ctn (red) or Ctn(15–34) (blue) at its MBC was added to 10^7 cfu/ml *E. coli* suspensions. Changes on the SYTOX® Green channel were monitored during 90 min immediately after the peptide addition. a, flow cytometry histograms on the SYTOX® Green channel at different time points in the presence of Ctn and Ctn(15–34). P_p is the percentage of permeabilization at each time interval (n) (in s). b, flow cytometry histograms on the SYTOX® Green channel for the negative (live bacteria) and positive (dead bacteria) controls, used to define gates. c, percentages of permeabilized bacteria over time after the addition of the peptides. Data were normalized by controls, corrected using the calibration curve shown in Fig. S8a, and fitted with the two-state model (34) as described under “Experimental procedures.” The kinetic curve for the untreated control shown in Fig. S8b demonstrates that bacteria remain alive during the whole acquisition, and the changes observed in c are due to the peptide activity.
Sensorgrams obtained with zwitterionic lipid systems (i.e. POPC and POPC/Chol/SM) (Fig. 5b, left) showed rapid dissociation, suggesting that Ctn(15–34) is rapidly removed from neutral membranes, and dose–response curves (Fig. 5b, right) showed that Ctn(15–34) has a weak overall affinity for these lipid membranes. No significant differences were detected between fluid or liquid-ordered phases, suggesting an overall weak affinity for neutral membranes. In contrast, Ctn(15–34) has higher affinity for vesicles made of anionic phospholipids, as suggested by dose-response curves and sensorgrams obtained with POPC/POPS and POPC/POPG (see Fig. 5b and Table 3). Improved affinity is observed with higher content of anionic lipids (e.g. compare POPC with POPC/POPS (80:20) and with POPC/POPS (60:40)). The addition of PE-phospholipids to anionic membranes did not improve Ctn(15–34) binding, as shown with POPC/POPS (80:20) versus POPC/POPE/POPG (60:20:20). Although an improvement in affinity was observed for all anionic membranes, Ctn(15–34) has better affinity for POPG than for POPC, as demonstrated by comparing its binding with POPC/POPG (60:40) and POPC/POPS (60:40) (Fig. 5b).

To investigate whether N-terminal RhB derivatization alters the binding behavior of Ctn(15–34), the labeled and unlabeled versions were compared for binding to POPC and POPC/POPS (80:20). Similar to the case for bacterial cells (Fig. 2 and Fig. 2, the RhB version shows higher affinity for both neutral and anionic membranes (Fig. S6).

Membrane-disrupting properties of Ctn(15–34)
To examine whether peptides disturb pure lipid bilayers, vesicles loaded with carboxyfluorescein (CF) were incubated with various concentrations of Ctn(15–34) or RhB-Ctn(15–34). Fig. 5c shows that Ctn(15–34) does not induce dye leakage from neutral POPC vesicles, yet it disrupts negatively charged POPC/POPG (80:20) vesicles in a dose-dependent manner (LC50 2.17 ± 0.36 μM). In agreement with an overall increase in affinity for the membrane, RhB-Ctn(15–34) disrupts vesicles, including zwitterionic membranes, with much higher effi-
Membrane-disruptive effect of Ctn and Ctn(15–34)

Figure 5. Interaction of Ctn(15–34) and RhB-Ctn(15–34) with LPS, LTA, model membranes, and RBCs. a, binding of Ctn(15–34) and of RhB-Ctn(15–34) to LTA and LPS as examined using the LAL assay. The concentration of RhB-Ctn(15–34) to neutralize 50% of LTA and LPS is 0.22 and 8.26 μM, respectively. Ctn(15–34) requires >32 μM to neutralize both LTA and LPS. b, binding of Ctn(15–34) to model membranes followed by surface plasmon resonance. Peptide samples were injected over lipid bilayer deposited onto an L1 chip surface; sensorgrams (left) obtained upon injection of 38 μM peptide over a particular lipid bilayer for 180 s (association phase) and the dissociation followed for 600 s (dissociation phase). Shown is the amount of Ctn(15–34) (right) obtained at the end of association phase (t = 170 s) upon injection of various peptide concentrations. The signal of sensorgrams and dose–response curves were normalized to peptide/lipid ratios (P/L, mol/mol) by converting response units into mol of peptide and normalized for the amount of lipid deposited onto the lipid surface (1 response unit = 1 pg/mm² of peptide or lipid). c, membrane disruption induced by Ctn(15–34) and RhB-Ctn(15–34). The percentage of membrane leakage was determined by CF dequenching. LUVs composed of POPC or POPC/POPS (80:20) with total lipid concentration of 5 mg/mL were incubated with various concentrations of peptide. Melittin, a membrane-disrupting peptide, was included as a positive control. d, toxicity against RBCs induced by Ctn(15–34) and RhB-Ctn(15–34). A suspension of RBCs (0.25%, v/v) was incubated with various concentrations of peptides. Percentage of hemolysis was detected by absorbance at 340 nm.

Table 3
Affinity of Ctn(15–34) for different lipid systems as followed by surface plasmon resonance

| Lipid system            | P/Lmax (mol/mol) |
|-------------------------|------------------|
| POPC                    | 0.33 ± 0.05      |
| POPC/Chol/SM (27:33:40) | 0.37 ± 0.02      |
| E. coli polar extract   | 0.77 ± 0.04      |
| POPC/POPS (80:20)       | 0.83 ± 0.11      |
| POPC/POPS (60:20)       | 1.76 ± 0.15      |
| POPC/POPS/POPE (60:20:20)| 0.65 ± 0.09    |
| POPC/POPG (80:20)       | 0.63 ± 0.07      |
| POPC/POPG (60:40)       | 1.35 ± 0.15      |

Toxicity to human red blood cells

The hemolytic properties of Ctn(15–34) and RhB-Ctn(15–34) were tested in human red blood cells (RBCs). Ctn(15–34) is not hemolytic up to 64 μM when tested against 0.25% (v/v) of RBCs (Fig. 5d). The lack of toxicity of Ctn(15–34) against RBCs is in agreement with its low affinity and inability to disturb neutral model membranes. On the other hand, derivatization of the peptide with RhB makes the peptide toxic, as shown with RhB-Ctn(15–34) (HC50 4.33 ± 0.18 μM) and is in agreement with this peptide possessing high affinity and the ability to disrupt lipid membranes.

Discussion

The aim of this work was to gain insights into the antimicrobial mechanisms of Ctn, an AMP derived from the venom gland of the rattlesnake Crotalus durissus terrificus, and of its fragment Ctn(15–34). Previous studies demonstrated that Ctn is toxic against bacteria, fungi, tumor, and healthy cells (26, 28). In contrast, Ctn(15–34) kills Gram-negative bacteria, fungi, and tumor cells but not healthy eukaryotic cells. Moreover, Ctn(15–34) has remarkable stability in human serum and is 40% smaller than Ctn (20 versus 34 amino acid residues) (27). For this reason, Ctn(15–34) is regarded as a promising anti-infective lead. However, its mechanism of action had not been examined until now.

To investigate the mode of action, we compared the antimicrobial potency and ability to disrupt membranes of Ctn and Ctn(15–34), using Gram-negative E. coli and P. aeruginosa as model organisms. We confirmed that both peptides are bactericidal at low micromolar concentrations (Table 1) and demonstrated that bacterial death is accomplished remarkably fast (Fig. 1a), particularly against P. aeruginosa. A fast induction of bacterial death is normally associated with loss of physical integrity of the membrane (39), as described previously for AMPs such as melittin (40), C16G2 (41), VAMP 059, and VCPP 2319 (42). The current results confirm a mode of action related to membrane disruption.
and suggest different stages in the permeabilization process (Fig. 6): (Stage 1) initial peptide recruitment; (Stage 2) peptide accumulation; and (Stage 3) cell death by membrane disruption.

**Stage 1: Initial peptide recruitment**

There is wide agreement that AMPs are usually drawn to microbial surfaces by electrostatic attraction between cationic (Lys, Arg, and His) residues in the peptide and anionic components in the bacterial surface, such as LPS and LTA on Gram-negative or Gram-positive surfaces, respectively, and/or anionic phospholipids (PG and cardiolipin) in the plasma membrane (7). In the present case, the high cationicity of both Ctn and Ctn(15–34) (net charge +16 and +8, respectively) and their preferential action on Gram-negative bacteria suggest that both peptides might bind LPS, as proposed for other snake-derived cathelicidins, such as Pb-CATH (43) and HC-CATH (22). As shown in Fig. 5a, Ctn(15–34) is not highly efficient at neutralizing both LPS and LTA, but even so, the binding is higher for LPS than for LTA, in concordance with its preference for Gram-negative bacteria. The fact that both peptides show a preference for cardiolipin (see below in AFM results) also reinforces an electrostatically driven initial recruitment step.

**Stage 2: Peptide accumulation**

Build-up of Ctn and Ctn(15–34) onto the bacterial surface was confirmed by ζ potential studies, because the negatively charged surface of both E. coli and P. aeruginosa was neutralized by increasing peptide concentrations, which caused a shift to positive ζ potential values (Fig. 1b). In all cases, neutralization occurred at concentrations lower than MIC or MBC (Fig. 1b), suggesting that surface charge equalization is not enough and that more peptide is necessary to induce bacterial death, which takes place after a threshold concentration of Ctn and Ctn(15–34) on the bacteria surface has been reached and peptides can begin to cross/enter the membrane.

**Stage 3: Cell death by membrane disruption**

Bacterial death results from membrane disruption and consequent loss of functionality. FCA data in Fig. 2 and Fig. S2 show a direct correlation between peptide uptake, membrane permeabilization, and viability loss. A mechanism dependent on the ability to target the membrane was also confirmed by imaging techniques. AFM studies showed important disturbances in the E. coli surface upon incubation with Ctn and Ctn(15–34) (Fig. 4b). The leakage of cytoplasmic content observed after peptide treatment is in agreement with a lytic mechanism of action, and the shrinkage observed in the middle region of the
bacteria could be due to preferential accumulation of peptides in cardiolipin-rich domains, namely in the apical and septal area of the cell (44), as described previously for other AMPs (32). Confocal microscopy experiments found RhB peptides in the periphery of E. coli bacteria (Fig. 4a), probably located on the membrane, in tune with the membrane-disruption hypothesis, although they were also found internalized inside cells in some cases. Thus, we propose that both peptides are initially recruited and accumulate around the surface until a threshold concentration is reached, after which they begin to enter/cross the lipid bilayer, causing cell death.

The membrane-lytic mechanism proposed here for Ctn and Ctn(15–34) is in agreement with results described for other snake-derived, evolutionarily related cathelicidins (e.g. cathelicidin-BF (Cbf), a 30-residue AMP from the elapid snake Bungarus fasciatus, also described as membrane-disruptive against bacteria) (45, 46). Analogs Cbf-K14 and Cbf-A11, able to kill multiresistant E. coli in a few hours, can also penetrate bacteria cell membrane as well as bind to DNA (47), suggesting that these peptides are not simply retained on the lipid bilayer. The N-terminal fragment, BF-15, and its analog ZY13, both active against resistant Candida albicans and less hemolytic than Cbf, also act in a membrane-disruptive manner (48). The aforementioned Pb-CATH and HC-CATH also possess bacterial membrane lytic properties (22, 43), and OH-CATH and NA-CATH, two cathelicidins found in cobra species, possess 79% identity with Ctn and are also membrane-active (24, 49).

Despite evidence suggesting a common membrane disruption mechanism for snake cathelicidins, how this process unfolds over time is not well studied, in part due to the limited information available by standard methods. To shed light on this issue, we employed time-resolved flow cytometry (34, 50), a high-resolution technique that allows monitoring peptide uptake and membrane permeabilization over time. This is particularly relevant to membrane-lytic peptides, for which binding and accumulation on the bacterial surface are early-onset events not easily monitored by end-point experiments. For instance, FCA kinetic curves (Fig. 3) reveal differences undetectable with the colony count method. Ctn(15–34) acts faster than Ctn due to faster binding and membrane permeabilization (Fig. 3). The slower stages in the action of Ctn cause a lag in its kinetic curve, although no cooperativity is detected, which demonstrates that common-sense reasoning that assigns lag phase directly to positive cooperativity (34, 51, 52) might be an oversimplification.

The mechanism of action of both Ctn and Ctn (15–34) includes steps whose kinetic curves deviate from the two-state model. Interestingly, the related NA-CATH cathelicidin seems to switch its mechanism of action from membrane disruption to pore-based lysis in a biphasic model, mimicking the E. coli membrane with an impact on kinetics similar to that described for Ctn (49). Given the high structural similarities between both peptides (an N-terminal α-helical segment followed by a disordered tail) (27, 53), it is plausible that Ctn may act similarly, and such behavior would explain the complex cell death kinetic pattern here reported. This dual behavior might be tentatively related to the composite structure (α-helix plus random tail) of both NA-CATH and Ctn, each moiety of the peptide embedding into the membrane in a different fashion, as suggested for NA-CATH (49). The fact that removal of the N-terminal helix of Ctn to give Ctn(15–34) results in a simplified killing mechanism corroborates our hypothesis.

RhB-labeled versions of Ctn and Ctn(15–34) show different uptake kinetics relative to unlabeled peptides (Fig. S4). RhB-Ctn binds slower than RhB-Ctn(15–34) (Fig. S4), but membrane permeabilization is much faster once the peptides are bound. Use of RhB-labeled Ctn(15–34) (Fig. S4) made possible the monitoring of peptide attachment to bacteria simultaneously with membrane permeabilization. RhB fluorophore emission confirms that RhB-Ctn(15–34) binds faster to membranes than RhB-Ctn, in agreement with the two-state model analyses for both peptides ($k_{on} = 7 \times 10^{-4}$ s$^{-1}$ versus $k_{off} = 2 \times 10^{-7}$ s$^{-1}$). In contrast, upon membrane binding, membrane permeabilization is faster for RhB-Ctn ($k_{on} = 1 \times 10^{-2}$ s$^{-1}$ versus $k_{off} = 1 \times 10^{-3}$ s$^{-1}$). For both labeled peptides, $f \sim 0$, which shows that no cooperativity exists. In general, kinetic and membrane-affinity results for RhB-Ctn(15–34) appear to be affected by RhB labeling (Fig. 5 (a, c, and d) and Figs. S2 and S6); hence, conclusions cannot be extrapolated to the unlabeled version.

The previously described putative selectivity of Ctn(15–34) for anionic domains (27) was corroborated by experiments with model membranes showing a preference for anionic, particularly PS-rich vesicles (Fig. 5b). LUV leakage (Fig. 5c) and RBC hemolysis (Fig. 5d) studies show that Ctn(15–34) preferentially interacts and disrupts vesicles mimicking bacterial and tumor rather than healthy eukaryotic cells. These results are in agreement with data from other snake cathelicidins (e.g. OH-CATH), which suggest a crucial implication of the four N-terminal amino acid residues in hemolytic activity (24).

In conclusion, by deploying a set of high-resolution methodologies, we have achieved accurate spatiotemporal characterization of events involved in the bactericidal action of Ctn and Ctn(15–34) on E. coli and P. aeruginosa. The study sheds light on the mechanism whereby the peptides accumulate and disrupt bacterial and model membranes and reveals slight differences in their behavior, including different kinetics of access to bacterial surfaces and different toxicity. By dissecting Ctn into its Ctn(15–34) fragment, we have produced a smaller, simpler, and more cost-efficient AMP displaying selectivity for bacterial-like, PS-rich membranes and a simplified mechanism of action. Taken together, our findings augur a promising role for Ctn(15–34) as an anti-infective lead and, in a broader sense, provide valuable insights into AMP lethal mechanisms that should be useful in the design and development of antimicrobials against resistant bacteria.

**Experimental procedures**

**Peptides**

Peptides in Table 1 were assembled as C-terminal carboxamides on Rink amide resin as described previously (27). For N-terminal RhB coupling, the peptide-resin was N-deprotected with piperidine/N,N-dimethylformamide (20:80, v/v) and treated with a 5-fold molar excess of RhB/N,N′-dipropylcarbodiimide (1:1 molar ratio) in dichloromethane, followed by extensive dichloromethane and N,N-dimethylformamide washes. Analytical
Membrane-disruptive effect of Ctn and Ctn(15–34)

reversed-phase HPLC, preparative HPLC, and LC-MS analyses as were reported previously (27). Peptide stock solutions were prepared in sterile Milli Q water and stored at −20 °C.

Antibacterial and bactericidal assays

The antimicrobial activity of Ctn and Ctn(15–34) was studied against Gram-negative reference strains E. coli ATCC 25922 and P. aeruginosa ATCC 27853. Minimal inhibitory and bactericidal concentrations (MIC and MBC) were determined by the broth microdilution method as described in document M26-A of the Clinical and Laboratory Standards Institute (54). Briefly, bacterial suspensions were prepared by the direct colony suspension method into BBL Mueller Hinton II Broth (cation-adjusted) medium (MHBII) (BD Biosciences) and incubated for 18 h at 37 °C with 2-fold serial peptide dilutions on 96-well microtiter polystyrene plates (Corning Inc.). Final bacterial concentrations were in the 0.1–100 μM range. After incubation, MIC values were determined as the lowest peptide concentration preventing visible bacterial growth. MBC values were subsequently determined as described previously (42). All assays were performed in triplicate.

Time-kill curves

The time-dependent bactericidal effect of Ctn and Ctn(15–34) were assessed against both E. coli and P. aeruginosa. Bacterial suspensions were treated with the peptides at a final concentration corresponding to their MBC and incubated at 37 °C and 200 rpm. As control, bacterial suspensions without peptide were prepared under the same conditions. Aliquots of untreated and peptide-treated bacterial suspensions were withdrawn at different time points (t = 5, 15, 30, 60, 90, 120, 180, and 240 min), serially diluted, plated onto trypticase soy agar plates, and incubated at 37 °C. After 24 h, the number of bacterial colonies was counted, and viable bacteria (in cfu/ml) at each t were calculated as percentage of the untreated control at t0. The assay was performed in triplicate.

ζ potential

For the ζ potential studies, E. coli or P. aeruginosa suspensions were prepared in HEPES buffer (10 mM HEPES, 150 mM NaCl, pH 7.4) at 107 cfu/ml, and Ctn or Ctn(15–34) was added at different concentrations. Treated and untreated suspensions were injected into zeta cells and allowed to equilibrate at 25 °C for 15 min, and measurements were acquired on a Zetasizer Nano ZS (Malvern Instruments, Worcestershire, UK). ζ potential was calculated from the mean of 15 measurements, using the Smoluchowski equation as described previously (55). Experiments were performed in triplicate.

Viability and membrane permeabilization correlation studies

To determine bacterial viability and membrane permeabilization, bacterial suspensions at 108 cfu/ml in MHBII were centrifuged 10 min at 4000 × g and diluted to 5 × 107 cfu/ml in HEPES buffer. Incubation with 2-fold serial peptide dilutions was done at 37 °C and 200 rpm, for 90 min for E. coli and 60 min for P. aeruginosa. Untreated bacteria were used as negative control. To estimate the percentage of permeabilized bacteria and percentage of viable bacteria, each sample was respectively evaluated by FCA and colony count methods using protocols described previously (42).

Membrane permeabilization induced by RhB-labeled peptides

Samples for FCA were prepared as described above. Data were acquired on a BD FACSaria III cell sorter (BD Biosciences) using blue (488-nm) and yellow-green (561-nm) lasers and emission filters BP530/30 (LP505) and BP586/15 to detect SYTOX® Green and RhB, respectively. 5000 events corresponding to bacterial population as previously defined by forward scatter and side scatter were recorded. Experiments were performed in triplicate.

Peptide uptake and bacterial death kinetics

Peptide uptake and bacterial permeabilization were monitored in real time as described elsewhere (34). Briefly, a 108 cfu/ml bacterial inoculum was prepared in MHBII, centrifuged for 10 min at 4000 × g, and diluted to 107 cfu/ml on HEPES buffer. Dead bacteria were prepared by incubation at 37 °C with 70% (v/v) isopropyl alcohol for 60 min. SYTOX® Green was added to the bacterial suspension at 0.05 μM final concentration and incubated for at least 10 min. Ctn, RhB-Ctn, Ctn(15–34), or RhB-Ctn(15–34) was added to a final concentration equal to its MBC. Measurements were performed on a Fortessa X-20 instrument (BD Biosciences) using blue (488 nm) and yellow-green (561 nm) lasers and emission filters BP530/30 (LP505) and BP586/15 to detect SYTOX® Green and RhB, respectively. A total of 10,000 events were recorded for controls. For kinetic experiments, acquisition was done for 90 min immediately after the peptide addition. A control of untreated E. coli was also recorded for 90 min to ensure that bacteria remained alive during the total acquisition (see Fig. S8b). The bacterial population was selected according to the forward scatter/side scatter and the positive/negative SYTOX® Green gates (SG+/SG−), and positive/negative RhB gates (RhB+/RhB−) were established using controls. For kinetic analysis, events (#) on the SG+ and SG− gates were clustered for time intervals (n) equal to 1 s, and the number of events for each interval (#,n) was counted. The ratios of permeabilized bacteria (P) and peptide uptake (1) (bound to the membrane or internalized) for each n were calculated as follows.

\[
P_n = \frac{#_n \text{ on SG}^+}{(#_n \text{ on SG}^+ + #_n \text{ on SG}^-)} \quad (\text{Eq. 1})
\]

\[
U_n = \frac{#_n \text{ on RhB}^+}{(#_n \text{ on RhB}^+ + #_n \text{ on RhB}^-)} \quad (\text{Eq. 2})
\]

Estimated percentages of permeabilized bacteria and peptide uptake for each n were calculated by the following equations.

\[
\% \text{ permeabilized bacteria at } n = \frac{P_n - P_{live}}{P_{dead} - P_{live}} \times 100 \quad (\text{Eq. 3})
\]

\[
\% \text{ peptide uptake at } n = \frac{U_n - U_{live}}{U_{dead} - U_{live}} \times 100 \quad (\text{Eq. 4})
\]

The percentage of permeabilized bacteria was corrected using the equation derived from the calibration curve shown on Fig. S8a, obtained by combining known percentages of dead and live bacteria.
Data were fitted to the two-state model described previously (34), using the following equation,

\[ y = y_0 + (B_l - y_0) \times \left( 1 + \frac{k'_1 e^{-k_y x} - k_2 e^{-k_x x}}{k_2 - k'_1} \right) \]  

(Eq. 5)

where \( k'_1 = k_{p,x}; y_0 = \) initial % permeabilized bacteria; \( B_l = \) final % permeabilized bacteria; \( k_2 = \) permeabilization kinetic constant; \( k_1 = \) membrane attachment constant; and \( f = \) cooperativity binding factor.

### Confocal microscopy

An *E. coli* suspension was prepared at 10⁸ cfu/ml in MHBII, centrifuged for 10 min at 4000 \( \times g \), and diluted to 10⁷ cfu/ml on HEPES buffer. RhB-labeled or unlabeled peptides were added to the bacterial suspensions at a final concentration equal to their MBC, and the samples were incubated at 37 °C and 200 rpm for 90 min. Untreated bacteria were used as a control. After incubation, samples were placed on ice, and SYTOX® Green was added at 5 µM final concentration. Samples were placed on an ibiTreat-coated 8-well µ-slide (Ibidi, Munich, Germany) before imaging. Acquisition was made on a confocal point-scanning Zeiss LSM 880 microscope (Carl Zeiss, Jena, Germany) equipped with an alpha Plan-ApoChromat ×100 oil immersion objective (1.46 numerical aperture). The 488-nm line from an argon laser and the DPSS 561-20 laser were used to excite SYTOX® Green and RhB, respectively. In the normal confocal mode, ×0.6 zoom images were recorded at 1024 × 1024 resolution. In the Airyscan superresolution mode, 5.5× zoom was applied to select one single bacteria cell, and the optimal resolution was selected. ZEN software was used for image acquisition and Airyscan image processing. Fiji software was used for background subtraction and brightness and contrast adjust (identically for compared image sets). At least 12 total images were acquired in three independent replicates.

### Atomic force microscopy

Morphological changes on *E. coli* cells induced by Ctn and Ctn(15–34) were visualized by AFM imaging. 10⁸ cfu/ml suspensions were prepared on MHBII, centrifuged for 10 min at 4000 \( \times g \), and diluted to 10⁷ cfu/ml in HEPES buffer. Bacterial suspensions in the absence of the presence of Ctn or Ctn(15–34) at their MBC were incubated at 37 °C and 200 rpm for 90 min. Untreated bacteria were used as a control. Slide preparation, image acquisition, and processing were as described previously (32).

### Binding affinity for LTA and LPS

Neutralization of LTA and LPS by Ctn(15–34) or RhB-Ctn(15–34) was examined using the end-point chromogenic LAL assay kit (QLC-1000, Lonza) following the same protocol as before (36).

### Vesicle leakage assay

Content leakage from POPC or POPC/POPS (80:20 molar ratio) vesicles was quantified by CF fluorescence dequenching using LUVs loaded with 50 mM CF as described previously (57). Briefly, 2-fold dilutions of peptides (starting with 10 µM) were incubated for 20 min with LUVs (5 µM of lipid concentration) in HEPES buffer. Fluorescence emission intensity (excitation at 494 nm and emission at 515 nm) was measured in an M1000 microplate reader (Tecan, Männedorf, Switzerland), and the leakage percentage was calculated as described previously (57).

### Interactions with lipid bilayers

Peptide-membrane interactions were monitored by SPR at 25 °C with a L1 biosensor chip in a Biacore 3000 instrument (GE Healthcare Australia, Parramatta, Australia). SUVs composed of POPC, POPC/POPE (80:20 molar ratio), POPC/POPG (80:20), POPC/POPG/Chol (27:33:40), POPC/POPS (80:20), POPC/POPS/POPE (60:40), and POPC/POPS/POPE (60:20:20), or *E. coli* polar extract were prepared using HEPES buffer and deposited onto a L1 chip by injecting SUV suspensions with 0.5 mM lipid at a flow rate of 2 µl/min for 40 min. Peptides were injected over the lipid bilayers at a flow rate of 5 µl/min for 180 s (association phase), and dissociation was monitored for 600 s (58, 59). HEPES was used as running buffer, and the chip was regenerated as before (60). All solutions were freshly prepared and filtered using a 0.22-µm pore size filter. The response units were normalized to peptide/lipid ratio as described previously (58).

### Hemolytic studies

Fresh human blood was collected from three healthy donors following protocols approved by the Human Research Ethics Unit at the University of Queensland. RBCs were isolated by centrifugation (1500 \( \times g \), 1 min), washed three times, and resuspended in PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 1.8 mM KH₂PO₄) at 0.25% (v/v). BRC suspensions were incubated with increasing concentrations of Ctn(15–34) or with RhB-Ctn(15–34) for 1 h at 37 °C, and the percentage of hemolysis was quantified by the 405-nm absorbance of released hemoglobin. Samples incubated with PBS or with Triton X-100 0.1% (v/v) were used to define 0 and 100% of hemolysis, respectively, as described previously (61). Melittin, a hemolytic peptide, was included as a control.

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