Peptide modification results in the formation of a dimer with a 60-fold enhanced antimicrobial activity

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Abstract

Cationic antimicrobial peptides (CAMPs) occur naturally in numerous organisms and are considered as a class of antibiotics with promising potential against multi-resistant bacteria. Herein, we report a strategy that can lead to the discovery of novel small CAMPs with greatly enhanced antimicrobial activity and retained antibiofilm potential. We geared our efforts towards \(i\) the N-terminal cysteine functionalization of a previously reported small synthetic cationic peptide (peptide 1037, KRFRIRVRV-NH\(_2\)), \(ii\) its dimerization through a disulfide bond, and \(iii\) a preliminary antimicrobial activity assessment of the newly prepared dimer against \textit{Pseudomonas aeruginosa} and \textit{Burkholderia cenocepacia}, pathogens responsible for the formation of biofilms in lungs of individuals with cystic fibrosis. This dimer is of high interest as it does not only show greatly enhanced bacterial growth inhibition properties compared to its pep1037 precursor (up to 60 times), but importantly, also displays antibiofilm potential at sub-MICs. Our results suggest that the reported dimer holds promise for its use in future adjunctive therapy, in combination with clinically-relevant antibiotics.

Introduction

Antibiotic resistance is a serious and growing phenomenon, as well as a primary public health concern [1, 2]. Unfortunately, antibiotics are no longer the magic bullets that they once were. New resistance mechanisms have emerged, making generations of antibiotics virtually ineffective, resulting in prolonged illness, greater risk of death and higher costs. Thus, development of new antibiotics and other novel strategies are critically needed. Biofilm-associated bacteria possess 10–1,000 fold greater resistance to antibiotic treatment compared to freely-floating, planktonic cells, making established biofilm infections especially difficult to eradicate [3, 4]. The biofilm mode of growth is a strategy used by microorganisms to survive unfavorable environmental conditions. An estimated 80% of all bacterial infections involve a biofilm component [5]. For instance, the severe antibiotic resistance of \textit{Pseudomonas aeruginosa} and \textit{Burkholderia cenocepacia} in lungs of Cystic Fibrosis (CF) patients has been associated with the formation of drug resistant biofilms [6, 7]. Bacteria benefit from the accumulation of thick and...
sticky mucus to colonize the lung tissue and airways in a multi-resistant biofilm, which results in respiratory failure and mortality [7]. To date, no antimicrobial that targets bacterial biofilm(s) has been clinically approved. Thus, there is a great interest in the development of antimicrobials that inhibit bacterial growth, and also prevent biofilm formation and/or promote established biofilm dissolution.

Cationic antimicrobial peptides (CAMPs) are a class of antibiotics with promising potential against multi-resistant bacteria. Various CAMPs with a broad spectrum activity were isolated from invertebrates, insects, microorganisms, plants, fish, amphibians, mammals and humans [8]. Attracted to the anionic phospholipid head, CAMPs establish strong nonspecific, hydrophobic and electrostatic interactions with the bacterial cytoplasmic membrane [9]. Although their therapeutic potential seems to be restricted, as it is the case for many other types of antimicrobials, by the difficulty to find a proper balance between critical parameters such as hydrophobicity, charge, length, amphiphilicity and structure, CAMPs have the advantage to be small and easily tunable. Still, their use remains restricted by their intrinsic characteristics, notably their toxicity to mammalian cells, limited tissue distribution and proteolytic degradation in the blood [10]. Among successful examples, the synthetic protegrin analog IB-367 is part of an aerosol formulation used after chemotherapy [11]. Although their mechanism of action is not completely established, it is recognized that CAMPs can interfere with different stages of bacterial biofilm formation. Initially, CAMPs prevent cell adhesion to the surface via electrostatic bonds with the membrane [12]. They are also capable of preventing biofilm maturation by acting on the molecular signals involved in quorum sensing [13]. A library of small synthetic CAMPs able to inhibit biofilm formation and promote biofilm degradation can be found in the BaAMPs database (biofilm active antimicrobial peptides database) [14].

The discovery of naturally occurring dimeric peptides with marked antimicrobial activity encouraged scientists to explore dimerization, including disulfide bridging and lactamization, as a means to confer to CAMPs many properties that may enhance their therapeutic potential as compared to their monomers [15, 16]. For instance, the Defr1 dimeric defensin peptide, containing a natural cysteine residue, has an activity against P. aeruginosa PAO1 (6 μg/mL) approximately 9-fold greater than its 34-residue monomer counterpart (50 μg/mL) [17]. Also, compared to its monomeric form, VG16KRKP dimer displays a 10-fold decrease of its MIC against E. coli and more than a 4-fold decrease against P. aeruginosa and K. pneumoniae [18]. The improved antibacterial activity was correlated with an increase in hydrophobicity and cationicity of its surface area, which enhances LPS binding and neutralization [16]. Interestingly, very few studies reported the effect of CAMP dimerization through cysteine disulfide bridge formation, especially for N-terminus cysteine-containing peptides [18].

Herein, we report the N-terminal functionalization of a small synthetic cationic peptide with a cysteine, its dimerization through a disulfide bond, and a preliminary antimicrobial activity assessment of the newly prepared dimer against two strains each of P. aeruginosa and B. cenocepacia. For this study, the synthetic 9-mer cationic peptide pep1037 (KRFRIRVRV-NH₂) was selected [19], due to its antimicrobial potential against P. aeruginosa and B. cenocepacia and its considerable antibiofilm activity. We report that such a strategy leads to the discovery of a small CAMP with greatly enhanced antimicrobial activity and retained antibiofilm potential.

Materials and methods
Peptide synthesis/modification
Pep1037 and cys-pep1037 were synthesized manually using a standard solid phase peptide synthesis approach with Fmoc chemistry (Fig 1). Couplings of the protected amino acids were
mediated by benzotriazol-1-yloxy-tris-(dimethylamino)phosphonium hexafluorophosphate (BOP; 3 equiv) and N,N’-diisopropylethylamine (DIPEA; 6 equiv) in N,N-dimethylformamide (DMF) at room temperature for 0.5–1 h. Coupling efficiency was monitored with the qualitative ninhydrin test and a 3-equivalent excess of the protected amino acids based on the original substitution of the Fmoc Rink-amide resin (0.62 mmol/g). Fmoc removal was achieved with 20% piperidine in DMF at room temperature for 10 minutes. Peptides were cleaved from the resin support with simultaneous side chain deprotection by treatment with TFA containing 1,2-ethanedithiol (2.5%), water (5%), triisopropylsilane (1%), thioanisole (5%) and phenol (5%) for 4 h at room temperature. After cleavage, the resin was removed by filtration, the filtrate was concentrated and peptides were precipitated using cold diethyl ether. Crude peptides were then solubilized in water and lyophilized prior to their purification using preparative RP-HPLC. Cys-pep1037 dimer was isolated by reversed phase HPLC from a lyophilized sample of crude cys-pep1037. Mal-cys-pep1037 was prepared and isolated using the following procedure: cys-pep1037 was dissolved to a final concentration of 1mg/mL in fresh PBS (phosphate buffer saline, pH = 7), and an excess of tris-carboxyethylphosphine (TCEP) reagent (10 equiv) was added (to prevent the formation of disulfide bonds). The mixture was stirred at room temperature for 20 min, an excess of maleimide (2 equiv) was added and was allowed to react for 4 h. Peptide mal-cys-pep1037 was isolated from the mixture by RP-HPLC.

All crude lyophilized peptides were purified using a preparative reversed-phase HPLC (RP-HPLC) protocol using a linear gradient from eluent A to B with 25% B per 2 min increments (Eluent A = H₂O, 0.1% TFA, Eluent B = CH₃CN/40% A, 0.1% TFA) and a C18 (column Phenomenex, Jupiter®), 15μm, 10 x 300 mm, 4 mL/min, absorbance measured at 220nm).
Homogeneity of purified fractions was assessed by RP-HPLC and MALDI-TOF mass spectrometry in linear mode using α-cyanohydroxycinnamic acid (matrix). Fractions containing the desired product were pooled and lyophilized. Overall, RP-HPLC analysis of each analog revealed that the purity of all peptides was ≥ 97%. For each peptide, the main peak observed by mass spectrometry (S1 Fig) was in agreement with the theoretical mass values ([M+H]+ m/z) 1229.3; cys-pep1037: m/z ([M+H]+) 1331.8; mal-cys-pep1037: m/z ([M+H]+) 1429.7; cys-pep1037 dimer: m/z ([M+H]+) 2662.5.

**Bacterial strains and growth conditions.**  *P. aeruginosa* ATCC 27853 (blood culture isolate, American Type Culture Collection, Manassas, USA), *P. aeruginosa* ATCC 15442 (animal room water bottle isolate), *B. cenocepacia* K56-2 (CF patient isolate) and *B. cenocepacia* J2315 (multidrug-resistant CF patient isolate) were used. Bacteria were subcultured on TSA (tryptic soy agar, BD Difco™ Dehydrated Culture Media, Fisher Scientific) plates from frozen stocks. Inocula was prepared in TSB (typtic soy broth, BD Difco™ Dehydrated Culture Media, Fisher scientific) in overnight cultures at 37°C with shaking at 200 rpm.

**Minimum Inhibitory Concentration (MIC) determination.** The broth microdilution method (with minor modifications for cationic peptides) was performed according to the guidelines of "Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically; Approved Standard-Ninth Edition" developed by the Clinical Laboratory Standards Institute’s committee [20]. Peptides were dissolved in water and stored in glass vials. Different peptide concentrations were prepared in 100 μL of cationic adjusted (25 mg/L CaCl₂ and 12.5 mg/L MgCl₂) MHB (Mueller Hinton broth, BD Difco™ Dehydrated Culture Media, Fisher Scientific) and added to sterile 96-well polypropylene microtiter plates. Each well was inoculated to a final concentration of 5 × 10⁶ CFU (50 μL of MHB inocula suspension prepared from colonies grown on MHA) [21]. Plates were incubated at 37°C under static conditions for 18 h and 24 h for *P. aeruginosa* and *B. cenocepacia*, respectively, and absorbance was read at 600 nm using a Cytation 3-cell imaging plate reader. The MIC was defined as the lowest concentration of peptide at which no growth was observed. All experiments were done at least in triplicate.

**Biofilm inhibition assay.** The ability of pep1037, cys-pep1037 dimer and mal-cys-pep1037 to prevent biofilm formation was evaluated by a method previously described [22, 23] with the following modifications. *P. aeruginosa* and *B. cenocepacia* cells (1 × 10⁵ CFU) were grown in 96-well polypropylene microtiter plates (BD falcon # 353912) in 200 μL final volume of TSB at 37°C for 24 h in the presence (2.5 μg/mL) or not of the peptides. Optical density (OD) of the cultures at 600 nm was determined after inoculation. As a measure of the planktonic bacterial growth, OD was again measured after 24 h of incubation, prior to biofilm staining. The planktonic suspension was gently aspirated using pipettes. Wells were then washed by 200 μL PBS and shaken out by turning the plate over. Biofilm cells were fixed by 200 μL of 99% methanol per well for 15 min. Then, methanol was aspirated and plates were air dried overnight. Biofilm production was measured using the crystal violet (CV, 1%) stain technique [23]. Briefly, excess CV was rinsed three times with water and plates were air-dried overnight. Stain was solubilized in 250 μL of 30% glacial acetic acid and quantified using a plate reader at 570 nm. All experiments were done in six replicates.

**Biofilm disintegration assay.** The pre-formed biofilm assay was performed as previously described [24]. *P. aeruginosa* ATCC 27853 or ATCC 15442 (1 × 10⁵ CFU) were incubated under static conditions in 200 μL of TSB for 24 h at 37°C. TSB media was then aspirated and the established biofilm was treated with various concentrations of pep1037, cys-pep1037 dimer and mal-cys-pep1037 solutions prepared in fresh TSB. The microtiter plate was incubated for 24 h, at 37°C. Biofilm production was measured using the crystal violet stain technique as described above. All experiments were done in six replicates.
Results and discussion

To assess the impact of pep1037 dimerization on bacterial growth and biofilm formation/degradation, a cysteine residue was introduced at its N-terminus, cys-pep1037. As reported for other cysteine containing peptides [25–28], dimerization occurs spontaneously during the lyophilization/purification step following the synthesis of cys-pep1037. Both peptides, cys-pep1037 and cys-pep1037 dimer, were isolated using RP-HPLC (Fig 1).

Inhibition of bacterial growth

The ability of cys-pep1037 dimer to inhibit the growth of P. aeruginosa was evaluated by determining its minimum inhibitory concentration (MIC) against two different strains of P. aeruginosa, notably ATCC 27853 and ATCC 15442, and results were compared to the MIC of native pep1037 against the same strains. MIC values determined for pep1037 were in agreement with previously reported MIC values of the same peptide against two Gram-negative pathogens, P. aeruginosa (PAO1 and PA14) and B. cenocepacia (4813), from 304 to > 608 μg/mL, respectively [19]. Interestingly, cys-pep1037 dimer was found to display a 30 to 60 times lower MIC than pep1037 against P. aeruginosa ATCC 27853 and ATCC 15442, respectively (Table 1). The MIC of pure cys-pep1037 could not be assessed due the presence of variable amounts of the corresponding dimer in the stock solutions used, but nevertheless, the MIC of this peptide mixture was found to be much lower than that of pep1037, but 4–8 times higher than that of cys-pep1037 dimer alone. This result strongly suggests that the antimicrobial activity of this peptide mixture is due to the presence of the dimer, although one cannot rule out the possible additional contribution of cys-pep1037. It is important to note that the dimer to monomer ratio might increase over time during MIC determination experiments, as considerable amounts of dimer were observed when solutions of cys-pep1037 (40 μg/mL) were incubated in culture medium for 1 h.

To prevent cys-pep1037 from undergoing dimerization, its thiol functionality was reacted with maleimide (Fig 2). Mal-cys-pep1037 displayed a lower antimicrobial activity against both strains of P. aeruginosa compared to its precursor, cys-pep1037, but an activity equivalent to that of pep1037. This again supports the previously mentioned hypothesis that the antimicrobial activity of the cys-pep1037 monomer/dimer mixture might be due to the presence of cys-pep1037 dimer, without completely ruling out an effect from the presence of an additional cysteine residue at the N-terminal position of pep1037. Indeed, bringing modifications at the N-terminal position of a peptide can have a drastic impact on its antimicrobial activity [29–31].

Table 1. Inhibitory effect of the peptides on bacterial growth\(^a\).

| Bacteria (strain)          | MIC (μg/mL) | pep1037\(^b\) | cys-pep1037\(^c\) | cys-pep1037 dimer | mal-cys-pep1037 |
|---------------------------|-------------|---------------|-------------------|-------------------|-----------------|
| P. aeruginosa (ATCC 27853)| 307         | 40            | 10                | 357               |
| P. aeruginosa (ATCC 15442)| 307         | 40            | 5                 | 357               |
| B. cenocepacia (J2315)    | > 614        | 332           | 332               | > 614             |
| B. cenocepacia (K56-2)    | > 614        | 332           | 332               | > 614             |

\(^a\)Identical results were obtained for all replicates.

\(^b\)Additional species and strains were also tested (see S1 Table).

\(^c\)Different stock solutions of cys-pep1037 (13 mg/mL) used for this experiment contained variable amounts of cys-pep1037 dimer.
All peptides were also tested against two strains of *B. cenocepacia*, bacteria that also play a role in CF lung infection. Their antibacterial potential against both strains was found to be at least two times higher than the one of pep1037 (332 μg/mL compared to > 614 μg/mL), but in contrast to what was noted for both *P. aeruginosa* strains, *cys-pep1037 dimer* did not show an enhancement in antibacterial activity compared to the *cys-pep1037* monomer / dimer mixture (Table 1). Dimerization of *cys-pep1037* from the monomer / dimer mixture in cultures could explain these results. This is further supported by the loss of *cys-pep1037* enhanced antibacterial effect (Table 1) upon reaction with maleimide, likely due to the inability of mal-cys-pep1037 to dimerize.

**Biofilm inhibition and disintegration**

To further explore the potential of *cys-pep1037 dimer*, we investigated whether the antibiofilm ability of its monomeric precursor pep1037 could be retained. The inhibitory effect of *cys-pep1037 dimer* on the formation of *P. aeruginosa* and *B. cenocepacia* biofilms was then assessed using the crystal violet assay (Figs 3 and 4). For comparison purposes, the inhibitory effect of mal-cys-pep1037 was also evaluated and results were compared to the ones obtained for pep1037 (which were found to be in agreement with previous reports) [19]. From these experiments, we conclude that *i*) the modification of pep1037 with a maleimide-protected cysteine at the N-terminal position (mal-cys-pep1037), and *ii*) the dimerization of its cysteine N-terminal adduct (cys-pep1037 dimer) do not lead to an overall loss in its ability to prevent biofilm formation. More specifically, at a concentration of 10 μg/mL, all peptides displayed a considerable effect on the formation of the four biofilms studied after 24h (Fig 3). A particularly pronounced effect of the peptides was noted against the *B. cenocepacia* J2315 biofilm (Fig 3D), 10 μg/mL being at least 30 times lower than their respective MIC (Table 1). Many CAMPs were reported to display a biofilm formation inhibitory effect at a concentration much lower than their MIC [29, 32, 33], but very few of these CAMPs were found to display a significant antimicrobial activity against planktonic bacteria [34]. Since the inhibitory effect observed for *cys-pep1037 dimer* on the formation of both *P. aeruginosa* strains could simply be due to its high potential to inhibit planktonic bacterial growth (MIC = 5–10, Table 1), the same experiment was also performed for all peptides at a concentration of 2.5 μg/mL. No significant inhibitory effect was observed on the formation of the four biofilms for any of the peptides when a sub-MIC concentration of 2.5 μg/mL was used (Fig 4) but interestingly, the combination of *cys-pep1037 dimer* (2.5 μg/mL) and *pep1037* (2.5 μg/mL) led to a considerable inhibitory effect in the range of 60–80% (Fig 4). This highlights the interest of further studying the potential use of sub-MIC concentrations of this dimer in adjunctive therapy.
The biofilm disintegration potential of pep1037, cys-pep1037 dimer and mal-cys-pep1037 was also assessed on a *P. aeruginosa* ATCC 15442 (A), *P. aeruginosa* ATCC 27853 (B), *B. cenocepa cia* K56-2 (C) and *B. cenocepa cia* J2315 (D) biofilms. All three peptides were found to have a considerable potential to disturb the established biofilm studied (Fig 5). A concentration of 10 μg/mL of pep1037 or mal-cys-pep1037 is sufficient to reduce by 56–60% the established monospecies biofilm, and there is no significant difference between the effect of both these peptides on biofilm degradation. Some CAMPs were previously reported to induce the dispersion of viable cells of mature biofilms [35]. Interestingly, for all concentrations studied, the established biofilm is reduced to a lesser extent by cys-pep1037 dimer than when treated with pep1037 or mal-cys-pep1037 (Fig 5). The effect of cys-pep1037 dimer on mature biofilm might not only be affected by its interaction with organisms in the *P. aeruginosa* biofilm but might also be strongly related to its ability to diffuse into the biofilm, as suggested for previously reported peptides [36]. The ability of cationic antibiotics to diffuse into biofilms is negatively influenced by an increase in molecular weight [37], and electrostatic interactions of cationic peptides with negatively charged biofilm matrix can delay CAMPs penetration [38]. These factors might explain why cys-pep1037 dimer is less active than its monomeric precursor pep1037 on the pre-formed biofilm tested. Nevertheless, the potential of the dimeric
form of cys-pep1037 to degrade an established P. aeruginosa biofilm is retained at sub-MIC concentrations.

**Conclusion**

We report here the N-terminal functionalization of a small synthetic cationic peptide with a cysteine, leading to the spontaneous formation of a dimer with a 30 to 60-fold enhancement in antimicrobial activity compared to its precursor. This work demonstrates that this strategy can potentially lead to the discovery of relatively small CAMPs with a greatly enhanced antimicrobial activity through a simple approach. Importantly, this can be achieved while retaining considerable activity against mature biofilms at sub-MICS. *P. aeruginosa* and *B. cenocepacia* are important pathogens responsible for various opportunistic infections, including the morbidity and mortality in CF patients via the formation of biofilms in lungs. These results suggest that cys-pep1037 dimer holds promise for its use in adjunctive therapy and future research may demonstrate potential for its use in combination with clinically-relevant antibiotics [39]. This study also opens the door to the improvement of small CAMPs via cysteine disulfide dimerization at the N-terminus, which has so far been unexplored.
Supporting information

S1 Table. Bacterial growth inhibitory effect of the peptides.
(DOCX)

S1 Fig. Mass spectral data for all peptides (MALDI-TOF mass spectrometry in linear mode using α-cyanohydroxycinnamic acid as matrix).
(DOCX)

Acknowledgments

The authors wish to thank Prof. Frédéric Veyrier for useful suggestions and Mrs. Marie-Christine Groleau for technical assistance.

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