# **Antimicrobial Peptides with Antiendotoxin**

Subjects: Microbiology Contributor: JYAWEI CHENG

A strategy was described to design antimicrobial peptides (AMPs) with enhanced salt resistance and antiendotoxin activities by linking two helical AMPs with the Ala-Gly-Pro (AGP) hinge. Among the designed peptides, KR12AGPWR6 demonstrated the best antimicrobial activities even in high salt conditions (NaCl ~300 mM) and possessed the strongest antiendotoxin activities. These activities may be related to hydrophobicity, membrane-permeability, and  $\alpha$ -helical content of the peptide. Amino acids of the C-terminal helices were found to affect the peptide-induced permeabilization of LUVs, the  $\alpha$ -helicity of the designed peptides under various LUVs, and the LPS aggregation and size alternation. A possible model was proposed to explain the mechanism of LPS neutralization by the designed peptides. These findings could provide a new approach for designing AMPs with enhanced salt resistance and antiendotoxin activities for potential therapeutic applications.

Keywords: antimicrobial peptide ; salt resistance ; lipopolysaccharide ; antiendotoxin ; cecropin-like

#### 1. Overview

Antimicrobial peptides (AMPs) have been found in the innate defense systems of plants, insects, and animals. AMPs can incorporate and disturb microbial membranes and hence cause their death. Owing to these reasons, AMPs may be the solution to the problem of bacterial resistance. Problems such as salt sensitivity, cost of synthesis, bioavailability, and stability have limited the therapeutic applications of antimicrobial peptides. Cecropins are a family of antimicrobial peptides widely found in the innate immune system of Cecropia moth. Cecropins exhibit broad spectra antimicrobial and anticancer activities <sup>[1]</sup>. The structures of cecropins are composed of 34–39 amino acids with an N-terminal amphipathic  $\alpha$ -helix, an AGP hinge and a hydrophobic C-terminal  $\alpha$ -helix. Recently, cecropin A and the cecropin-like peptide papiliocin were found to possess anti-inflammatory activities in LPS-stimulated murine macrophage. Studies of cecropin analogues, cecropin A/cecropin B hybrids, and cecropin A, LL-37, and magainin hybrids also revealed potential antimicrobial and anticancer activities. We hypothesize that the binding and neutralization of LPS of cecropin and cecropin-like peptides is through similar structural features like S1-Nal-Nal <sup>[2]</sup>. Here, we use this structural feature to create a new type of antimicrobial peptides by linking an amphipathic peptide and a hydrophobic peptide with the AGP sequence. In the study, we demonstrated the antimicrobial and LPS neutralization activities of these designed peptides.

# 2. Antimicrobial Peptides

To investigate the amphipathic helix–AGP–hydrophobic helix on the antimicrobial and neutralization activities of LPS, two peptides KR12 and RW6 were used for the N-terminal and C-terminal helices. KR-12 (KRIVQRIKDFLR) was derived from human host defense cathelicidin LL-37 and was found to possess antimicrobial activities against Gram-negative and Gram-positive bacteria. RW6 (RRWWRW) was derived from the reversed sequence of WR6 (WRWWRR) which had moderate antimicrobial activities. The helical wheel analyses of the designed peptides are shown in **Figure 1**. KR12AGPKR6, KR12AGPWR6 and KR12AGPVR6 were designed to compare the effects of hydrophobicity, membrane-permeability, and  $\alpha$ -helical contents. All peptides were acetylated and amidated at the N- and C-terminus.



Figure 1. Helical wheel of the peptides. (A) cecropin A, (B) cecropin B, (C) KR12, (D) RW6, (E) KR12AGPKR6, (F) KR12AGPWR6, (G) KR12AGPVR6.

#### 3. Antibacterial Activity and Salt Resistance

Minimal inhibitory concentration values (MICs) of the peptides were determined against Gram-positive and Gramnegative bacteria in Mueller–Hinton (MH) broth or LYM broth media (with 50, 100, 200 or 300 mM NaCl added) (**Figure 2**). KR12 was very effective against both Gram-positive and negative bacteria in LYM medium (MIC ~1 µg/mL). However, MIC values of KR12 increased to 4~16 µg/mL in MH broth and ~32 µg/mL in LYM broth with 300mM NaCl added. RW6 had only limited antimicrobial activity (8~32 µg/mL) in LYM broth and lost its antibacterial activity in MH broth and in LYM broth with >100 mM NaCl added. Similar to KR12, KR12AGPKR6 had MICs 1~2 µg/mL against both Gram-positive and negative bacteria in LYM medium. However, KR12AGPKR6 lost its antibacterial activity gradually to ~32 µg/mL in MH broth. KR12AGPKR6 was completely inactive in LYM broth with 300 mM NaCl added for *Escherichia coli*, *Staphylococcus aureus*, and *Pseudomonas aerginosa Migula*. Surprisingly, KR12AGPKR6 still possessed effective antibacterial activity against *Acinetobacter sp.* In MH broth (~2 µg/mL) and in LYM broth with 300 mM NaCl added (~1 µg/mL). On the other hand, KR12AGPWR6 displayed superior antibacterial activities against both Gram-positive and Gram-negative bacteria in MH and LYM media even in NaCl ~300 mM (MIC 2~4 µg/mL). KR12AGPVR6 exhibited moderate activities in MH broth (4~32 µg/mL), and lost its activity in LYM broth with 300 mM NaCl added. In addition, KR12, KR12AGPKR6, KR12AGPWR6 and KR12AGPVR6 were more effective to inhibit *Acinetobacter* sp. than the other three bacteria strains.



Figure 2. Minimal inhibitory concentration (MIC) values were displayed on a color scale for KR12, RW6, KR12AGPKR6, KR12AGPWR6 and KR12AGPVR6 under Mueller–Hinton (MH) broth and LYM medium with different concentrations of NaCl.

#### 4. Limulus Amebocyte Lysate (LAL) Assay

The ability of binding and neutralizing LPS in vitro was measured by the LAL assay which is well-known as the most sensitive and specific method. As shown in **Figure 3**, KR12AGPWR6 blocked the interaction between LPS and factor C (an LPS sensitive serine protease isolated from the hemocyte granules of the horseshoe crab Limulus), and neutralized the downstream reaction dose-dependently (greater than 60% inhibition at 64 µg/mL). KR12, RW6, KR12AGPKR6 and KR12AGPVR6 had only limited effects (less than 25% inhibition at 64 µg/mL).



**Figure 3.** Lipopolysaccharide (LPS) -neutralizing activities determined by *limulus* amebocyte lysate (LAL) assay. The experiments were performed in triplicate. Results are presented as means ± standard deviations (SD).

### 5. Endotoxemia Mouse Model

To evaluate the anti-LPS effect of KR12AGPWR6 in vivo, the mice were divided into three groups and received PBS, LPS, and a mixture of LPS and peptide, respectively. Blood was collected via the tail veins of the mice 1.5 h after injection. Serum TNF- $\alpha$  production increased dramatically in the LPS group. The KR12AGPWR6-treated mice displayed a significantly lower level of TNF- $\alpha$  (**Figure** 4A). Pathological evaluation of the lung tissues revealed that the proliferation of alveolar epithelial cells and pulmonary hemorrhage were reduced in mice treated with KR12AGPWR6 (**Figure** 4B–D).



**Figure 4.** Suppression of LPS-stimulated inflammation in endotoxemic mice (C57BL/6) by KR12AGPWR6. (A) LPSstimulated TNF- $\alpha$  release 1.5 h after injection. Excised lungs from the sacrificed mice 24 h after injection were subjected to H&E staining (B) control, (C) LPS (D) LPS and peptide. KR12AGPWR6 displayed the protective activity on LPSstimulated endotoxemic mice. All scale bars = 50 µm. Results are presented as means ± standard deviations (SD); *n* = 5. \* *p* < 0.05 versus LPS.

# 6. LPS Aggregation

It has been reported that LPS aggregation promoted by polymyxin B and rBPI<sub>21</sub> might inhibit the interaction of LPS with its cell receptors and hence block cytokine production. Dynamic light scattering (DLS) was used to measure the size increase of the designed peptides. The results indicated that KR12AGPWR6 promoted POPC/LPS LUVs aggregation and increased their mean sizes. KR12AGPKR6 and KR12AGPVR6 did not have the abilities in aggregating and increasing particle size (Figure 5).



**Figure 5.** Size distribution of LPS aggregates in the presence of the designed peptides. Plots showed the size alterations of LPS aggregates in the absence or presence of KR12AGPKR6, KR12AGPWR6 and KR12AGPVR6 in 25  $\mu$ M POPC/LPS solutions.

#### 7. Discussion

In summary, we describe a strategy to design AMPs with enhanced salt resistance and antiendotoxin activities by linking two helical AMPs with the AGP hinge. Among the designed peptides, KR12AGPWR6 demonstrated the best antibacterial activities even in high salt conditions (NaCl ~300 mM) and possessed the strongest antiendotoxin activities. These activities may be related to hydrophobicity, membrane-permeability, and  $\alpha$ -helical content of the peptide. Our results provide a new approach to design and development of AMPs with antimicrobial and antiendotoxin activities for potential therapeutic applications.



**Figure 6.** A proposed model of interaction of the designed peptide with LPS. The N-terminal helix of designed peptides shown in red and C-terminal hydrophobic helix shown in light blue. Structure of LPS consists three regions: polysaccharide O antigen (purple), core oligosaccharide (blue presents outer core and green presents inner core) and lipid A region. Lipid A consists of two glucosamine (yellow) units with attached acyl chains.

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