Research Article

Comparison of Biophysical and Biologic Properties of α-Helical Enantiomeric Antimicrobial Peptides

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In our previous study (Chen et al. J Biol Chem 2005, 280:12316–12329), we utilized an α -helical antimicrobial peptide V₆₈₁ as the framework to study the effects of peptide hydrophobicity, amphipathicity, and helicity on biologic activities where we obtained several $V_{\rm 681}$ analogs with dramatic improvement in peptide therapeutic indices against gram-negative and gram-positive bacteria. In the present study, the **D**-enantiomers of three peptides - V₆₈₁, V13A_D and V13K_L were synthesized to compare biophysical and biologic properties with their enantiomeric isomers. Each D-enantiomer was shown by circular dichroism spectroscopy to be a mirror image of the corresponding L-isomer in benign conditions and in the presence of 50% trifluoroethanol. L- and D-enantiomers exhibited equivalent antimicrobial activities against a diverse group of Pseudomonas aeruginosa clinical isolates, various gram-negative and gram-positive bacteria and a fungus. In addition, L- and D-enantiomeric peptides were equally active in their ability to lyse human red blood cells. The similar activity of L- and **D-enantiomeric peptides on prokaryotic or eukary**otic cell membranes suggests that there are no chiral receptors and the cell membrane is the sole target for these peptides. Peptide D-V13KD showed significant improvements in the therapeutic indices compared with the parent peptide V₆₈₁ by 53-fold against P. aeruginosa strains, 80-fold against gramnegative bacteria, 69-fold against gram-positive bacteria, and 33-fold against Candida albicans. The excellent stability of D-enantiomers to trypsin digestion (no proteolysis by trypsin) compared with the rapid breakdown of the L-enantiomers highlights the advantage of the D-enantiomers and their potential as clinical therapeutics.

Key words: α-helical peptides, antimicrobial activity, antimicrobial peptides, enantiomers, hemolytic activity, mechanism

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The widespread use of traditional antibiotics has resulted in the emergence of many antibiotic-resistant strains, prompting an urgent need for a new class of antibiotics (1,2). Cationic antimicrobial peptides have become important candidates as potential therapeutic agents (3-5) and have been shown to be active in in vivo animal studies (6). Although the exact mode of action of antimicrobial peptides has not been established (7-9), it has been proposed that the cytoplasmic membrane is the main target for many of these peptides, whereby peptide accumulation in the membrane causes increased permeability and loss of barrier function (10,11). The development of resistance to membrane active peptides whose sole target is the cytoplasmic membrane is not expected because this would require substantial changes in the lipid composition of cell membranes of micro-organisms. However, the major barrier to the use of antimicrobial peptides as antibiotics is their toxicity or ability to lyse eukaryotic cells, normally expressed as hemolytic activity (toxicity to human red blood cells), which is a main reason preventing their applications as injectable therapeutics.

Enantiomeric forms of antimicrobial peptides with all-D-amino acids were used to study the membrane-binding mechanism (12–14), as it was previously thought that cell membrane chirality would require a specific peptide chirality for it to be active. However, many studies have shown that all-D-amino acid peptides have equal activities to their all-L-enantiomers (12–19), suggesting that the antimicrobial mechanism of these peptides does not involve a stereoselective interaction with a chiral enzyme or lipid or protein receptor. In addition, all-D-peptides are resistant to proteolytic enzyme degradation, which enhances their potential as clinical therapeutics.

In our previous study (20), we utilized a *de novo* design approach to alter the secondary structure, hydrophobicity, and amphipathicity of an α -helical amphipathic antimicrobial peptide V₆₈₁ (21,22) and obtained lead compounds with high antimicrobial activities and

extremely low hemolytic activity. In the present study, we compare the biophysical and biologic properties between peptide enantiomers with all-L- or all-D-amino acids of our lead compounds, including antimicrobial activities against various bacterial strains, especially a diverse group of Pseudomonas aeruginosa clinical isolates with a wide range of susceptibility to the antibiotic ciprofloxacin. It is well known that respiratory infections by P. aeruginosa are the major cause of morbidity and mortality in adult patients with cystic fibrosis (23-25). Pseudomonas aeruginosa infection is also a serious problem in patients hospitalized with cancer and burns, the case fatality in such patients being 50% (24,25). According to data collected from 1990 to 1996 by the US Centers for Disease Control and Prevention (CDC), P. aeruginosa was the second most common cause of nosocomial pneumonia (17% of isolates), the third most common cause of urinary tract infections (11%), the fourth most common cause of surgical site infections (8%), the seventh most common isolated pathogen from the bloodstream (3%), and the fifth most common isolate overall (9%). We believe that, this comparative de novo design study of L- and D-antimicrobial peptides is the critical step toward the development of new antimicrobial therapeutics and understanding the mechanism of action of α -helical antimicrobial peptides.

Materials and Methods

Peptide synthesis and purification

Syntheses of the peptides were carried out by solid-phase peptide synthesis using *t*-butyloxycarbonyl chemistry and 4-methylbenzhydrylamine (MBHA) resin (0.97 mmol/g), as described previously (26). The crude peptides were purified by preparative reversed-phase high-performance liquid chromatography (RP-HPLC) using a Zorbax 300 SB-C₈ column (250 × 9.4 mm I.D.; 6.5 μ m particle size, 300 Å pore size; Agilent Technologies, Little Falls, DE, USA) with a linear AB gradient (0.1% acetonitrile/min) at a flow rate of 2 mL/min, where eluent A was 0.2% aqueous trifluoroacetic acid (TFA), pH 2 and eluent B was 0.2% TFA in acetonitrile (27). The purity of peptides was verified by analytical RP-HPLC as described below. The peptides were further characterized by mass spectrometry and amino acid analysis.

Analytical RP-HPLC of peptides

Peptides were analyzed on an Agilent 1100 series liquid chromatograph (Little Falls, DE, USA). Runs were performed on a Zorbax 300 SB-C₈ column (150 × 2.1 mm l.D.; 5 μ m particle size, 300 Å pore size) from Agilent Technologies using a linear AB gradient (1% acetonitrile/min) and a flow rate of 0.25 mL/min, where eluent A was 0.2% aqueous TFA, pH 2 and eluent B was 0.2% TFA in acetonitrile. Temperature profiling analyses were performed in 3 °C increments, from 5 to 80 °C using a linear AB gradient of 0.5% acetonitrile/min.

Characterization of helical structure

The mean residue molar ellipticities of peptides were determined by circular dichroism (CD) spectroscopy, using a Jasco J-720 spectropolarimeter (Easton, MD, USA), at 5 °C under benign (non-denaturing) conditions (50 mm KH₂PO₄/K₂HPO₄/100 mm KCI, pH 7.4), hereafter referred to as KP buffer, as well as in the presence of an α -helix inducing solvent, 2,2,2-trifluoroethanol (TFE; 50 mm KH₂PO₄/K₂HPO₄/100 mm KC1, pH 7.4 buffer/50% TFE). A 10-fold dilution of an approximately 500 μ M stock solution of the peptide analogs was loaded into a 0.02 cm quartz cell and its ellipticity scanned from 190 to 250 nm.

Measurement of antimicrobial activity (MIC)

Minimal inhibitory concentrations (MIC) were determined using a standard microtiter dilution method in a Mueller-Hinton (MH) medium. Briefly, cells were grown overnight

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at 37 °C in MH broth and diluted in the same medium. Serial dilutions of the peptides were added to the microtiter plates in a volume of 100 μ L followed by 10 μ L of bacteria to give a final inoculum of 1 × 10⁵ colony-forming units (CFU)/mL. Plates were incubated at 37 °C for 24 h and MICs determined as the lowest peptide concentration that inhibited growth. However, for MIC determination of *P. aeruginosa* clinical isolates, brain heart infusion (BH1) medium was used instead of MH broth. In addition, the bacteria were diluted to a final inoculum of 1 × 10⁶ CFU/mL.

Measurement of hemolytic activity (MHC)

Peptide samples were added to 1% human erythrocytes in phosphate-buffered saline (100 mM NaCl; 80 mM Na₂HPO₄; 20 mM NaH₂PO₄, pH 7.4) and reactions were incubated at 37 °C for 18 h in microtiter plates. Peptide samples were diluted twofold in order to determine the concentration that produced no hemolysis. This determination was made by withdrawing aliquots from the hemolysis assays, removing unlysed erythrocytes by centrifugation (800 × g) and determining which concentration of peptide failed to cause the release of hemoglobin. Hemoglobin release was determined spectrophotometrically at 570 nm. The hemolytic titre was the highest twofold dilution of the peptide that still cause release of hemoglobin from erythrocytes. The control for no release of hemoglobin was a sample of 1% erythrocytes without any peptide added. As erythrocytes were in an isotonic medium, no detectable release (<1% of that released upon complete hemolysis) of hemoglobin was observed from this control during the course of the assay. For the hemolysis time study, hemolytic activity of peptides at concentrations of 8, 16, 32, 64, 125, 250 and 500 μ g/mL was measured at 0, 1, 2, 4 and 8 h at 37 °C.

Calculation of therapeutic index (MHC/MIC ratio)

It should be noted that both MHC and MIC values were determined by serial twofold dilutions. Thus, for individual bacteria and individual peptides the therapeutic index (MHC/MIC) could vary by as much as fourfold if the peptide is very active in both hemolytic and antimicrobial activities; of course, if a peptide has poor or no hemolytic activity, the major variation in the therapeutic index (MHC/MIC) comes from the variation in the MIC value (as much as twofold).

Proteolytic stability assay

Proteolytic stability of the peptides was carried out with trypsin in a molar ratio of 1:20 000 (trypsin:peptide = 0.1 μ M:2 mM). The buffer used was 50 mM NH₄HCO₃ at pH 7.4 for both peptides and enzyme. The mixtures of peptide and trypsin were incubated at 37 °C. Samples were collected at time-points of 0, 5, 10, 20 and 30 min, 1, 2, 4 and 8 h. Equal volumes of 20% aqueous TFA were added to each sample to stop the reaction and peptide degradation was checked by RP-HPLC. Runs were performed on a Zorbax 300 SB-C₈ column (150 × 2.1 mm I.D.; 5 μ m particle size, 300 Å pore size) from Agilent Technologies at room temperature using a linear AB gradient (1% aceto-nitrile/min) and a flow rate of 0.25 mL/min, where eluent A was 0.2% aqueous TFA, pH 2 and eluent B was 0.2% TFA in acetonitrile. The change in integrated peak area of the peptides was used to monitor the degree of proteolysis during the time study.

Results

Peptide design

Peptide V₆₈₁ is a 26-residue α -helical amphipathic antimicrobial peptide (Figure 1 and Table 1) with excellent antimicrobial activity but high toxicity to human cells. Peptides V13K_L and V13A_D are peptides with L-Lys and D-Ala substitutions at position 13 of V₆₈₁ on the non-polar face of the helix (Table 1) and were chosen as lead peptide analogs for the present study because they demonstrated the best overall therapeutic indices against gram-negative and gram-positive bacteria, and the lowest toxicity against eukaryotic cells from a previous study (20). In the present study, we prepared the enantiomeric peptides of V₆₈₁ and analogs V13K_L and V13A_D.

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Table 1: Peptides used in this study

Name ^a	Peptide sequence ^b		
	1	13	26
V ₆₈	$AC - K_L - W_L - K_L$	$-S_{L}-F_{L}-L_{L}-K_{L}-T_{L}-F_{L}-K_{L}-S_{L}-A_{L}-V_{L}-K_{L}-T_{L}-V_{L}-L_{L}-H_{L}-H_{L}-A_{L}-L_{L}-K_{L}-A_{L}-L_{L}-S_{$	-S _L -amide
V13K	$_{\rm L}$ Ac $-$ K _L $-$ W _L $-$ K _L	$ = \sum_{L} - \sum$	$-S_{L}$ -amide
D-V13K V13A	$ \begin{array}{c} {}_{\mathrm{D}} \operatorname{A} \operatorname{C} - \operatorname{K}_{\mathrm{D}} - \operatorname{W}_{\mathrm{D}} - \operatorname{K}_{\mathrm{D}} \\ {}_{\mathrm{D}} \operatorname{A} \operatorname{C} - \operatorname{K}_{\mathrm{L}} - \operatorname{W}_{\mathrm{L}} - \operatorname{K}_{\mathrm{L}} \end{array} $	$-\mathbf{S}_{\mathrm{D}}-\mathbf{F}_{\mathrm{D}}-\mathbf{L}_{\mathrm{D}}-\mathbf{K}_{\mathrm{D}}-\mathbf{T}_{\mathrm{D}}-\mathbf{F}_{\mathrm{D}}-\mathbf{K}_{\mathrm{D}}-\mathbf{S}_{\mathrm{D}}-\mathbf{A}_{\mathrm{D}}-\mathbf{K}_{\mathrm{D}}-\mathbf{K}_{\mathrm{D}}-\mathbf{V}_{\mathrm{D}}-\mathbf{L}_{\mathrm{D}}-\mathbf{H}_{\mathrm{D}}-\mathbf{T}_{\mathrm{D}}-\mathbf{A}_{\mathrm{D}}-\mathbf{L}_{\mathrm{D}}-\mathbf{A}_{\mathrm{D}}-\mathbf{I}_{\mathrm{D}}-\mathbf{A}_{\mathrm{D}}-\mathbf{I}_{\mathrm{D}}-\mathbf{S}_{\mathrm{D}}\\ -\mathbf{S}_{\mathrm{L}}-\mathbf{F}_{\mathrm{L}}-\mathbf{L}_{\mathrm{L}}-\mathbf{K}_{\mathrm{L}}-\mathbf{T}_{\mathrm{L}}-\mathbf{K}_{\mathrm{L}}-\mathbf{S}_{\mathrm{L}}-\mathbf{A}_{\mathrm{D}}-\mathbf{L}-\mathbf{L}_{\mathrm{D}}-\mathbf{L}-\mathbf{L}-\mathbf{L}-\mathbf{L}-\mathbf{L}-\mathbf{L}-\mathbf{L}-$	$-S_{D}$ -amide $-S_{L}$ -amide
D-V13A C	$^{L} AC - K_{D} - W_{D} - K_{D}$ $^{C} AC - E_{L} - L_{L} - E_{L}$	$\begin{array}{l} -S_{\rm D} - F_{\rm D} - L_{\rm D} - K_{\rm D} - T_{\rm D} - F_{\rm D} - K_{\rm D} - S_{\rm D} - A_{\rm D} - A_{\rm L} - K_{\rm D} - T_{\rm D} - V_{\rm D} - L_{\rm D} - H_{\rm D} - T_{\rm D} - A_{\rm D} - L_{\rm D} - K_{\rm D} - A_{\rm D} - I_{\rm D} - S_{\rm D} \\ -K_{\rm L} - G_{\rm L} - G_{\rm L} - G_{\rm L} - E_{\rm L} - K_{\rm L} - G_{\rm L} - K_{\rm L} - E_{\rm L} - L_{\rm L} - E_{\rm L} - K_{\rm L} - amide \end{array}$	$-S_{D}$ -amide

^aD- denotes that all amino acids in the peptide are D-amino acids except position 13, which is denoted as L or D depending on the peptide.

^bPeptide sequences are shown using the one-letter code for amino acid residues; subscripts of L or D denote the L-amino acid and D-amino acid, respectively; Ac- denotes N^{α} -acetyl and -amide denotes C-terminal amide; in sequences, bold letters denote the amino acid substitutions at position 13 of the non-polar face of V_{R81} or D-V_{R81}.

^cPeptide C is a random coil peptide used as a control in reversed-phase high-performance liquid chromatography (RP-HPLC) temperature profiling.

Peptides V₆₈₁ and V13K_L contain all-L-amino acids and D-V₆₈₁ and D-V13K_D contain all-D-amino acids. In the case of V13A_D and D-V13A_L, position 13 is D-alanine and L-alanine, respectively (Table 1). Thus, D-V₆₈₁, D-V13K_D, and D-V13A_L are opposite in stereochemistry to the corresponding L-peptides, V₆₈₁, V13K_L and V13A_D, respectively. A control peptide C designed to exhibit negligible secondary structure, i.e. a random coil, was employed as a standard peptide for temperature profiling during RP-HPLC to monitor peptide association (Table 1) (20,28,29).

Secondary structure of peptides

To determine the secondary structure of peptides in different environments, CD spectra of the peptide analogs were measured under benign conditions (100 mm KCl, 50 mm KH_2PO_4/K_2HPO_4 , pH 7.4, referred to as KP buffer) and also in 50% TFE to mimic the hydrophobic environment of the membrane. As illustrated in Figure 2, the

parent peptide, V₆₈₁, was only partially helical in KP buffer; peptides V13K_L and V13A_D exhibited negligible secondary structure in KP buffer due to disruption of the non-polar face of the helix by introducing a hydrophilic L-lysine residue into peptide V13K_L or a helix-disruptive D-alanine residue into peptide V13A_D. However, in the presence of 50% TFE, all three L-peptides were fully folded α -helical structures with similar ellipticities and helicity (Table 2). As expected, the D-peptides showed spectra that were exact mirror images compared to their L-enantiomers, with ellipticities equivalent but of opposite sign both in benign KP buffer and in 50% TFE (Table 2).

Peptide self-association

The ability of the peptides to self-associate was determined by RP-HPLC temperature profiling. Figure 3A shows the change in RP-HPLC retention times of the three pairs of enantiomers and the



Figure 2: Circular dichroism (CD) spectra of peptides V₆₈₁ and D-V₆₈₁ (panel A), V13K_L and D-V13K_D (panel B) and peptides V13A_D and D-V13A_L (panel C) at pH 7.4 and 5 °C, in 50 mm aq. potassium phosphate buffer (KP buffer) containing 100 mM KCI. In panels (A-C), solid symbols represent the CD spectra of peptide analogs in KP buffer without trifluoroethanol (TFE), whilst open symbols represent CD spectra obtained in the presence of 50% TFE; the symbols used are: circles for L-peptides $V_{681},\ V13K_L,\ and\ V13A_D$ and diamonds for D-peptides D-V₆₈₁, D-V13K_D, and D-V13A_L.

control peptide C over a temperature range of 5–80 °C. As expected, L- and D-peptide enantiomers were totally inseparable over this temperature range, because each pair of peptides is identical in sequence and must adopt identical conformations on interacting with the RP matrix, whether in an all-L- or all-D-conformation. RP-HPLC retention behavior has been frequently utilized to represent overall peptide hydrophobicity (20,26,30,31). In the present study, the hydrophobicity of the three peptide pairs is in the order V₆₈₁/D-V₆₈₁ > V13A_D/D-V13A_L > V13K_L/D-V13K_D (Table 2), which agrees with the change in hydrophobic to the least hydrophobic amino acid residue (Val in V681 > Ala in V13A > Lys in V13K) (32). Figure 3B shows the retention behavior of the peptides after

normalization of the retention times to the corresponding retention time at 5 °C in order to highlight differences in the elution behavior of peptides as the temperature is increased from 5 to 80 °C. For example, the retention times of peptides V₆₈₁/D-V₆₈₁ increase with increasing temperature (up to approximately 30 °C) followed by a retention time decrease with a further temperature increase. Such a temperature profile is characteristic of a peptide exhibiting selfassociation (20,28,29,33). As illustrated in Figure 3C, the peptide self-association parameter, P_{A} , represents the maximum change in peptide retention time relative to the random coil peptide C. As peptide C is a monomeric random coil peptide in both aqueous and hydrophobic media, its retention behavior over the temperature ranging from 5 to 80 °C represents only general temperature effects

	Hydrophobi	city ^b	Benign		50% TFE		
Peptide ^a	t ⁵ _R (min)	t_R⁸⁰ (min)	[θ] ₂₂₂ ^c	% helix ^d	[θ] ₂₂₂ ^c	% helix ^d	PAe
V ₆₈₁	96.6	88.8	-13 000	46	-27 950	99	7.2
D-V ₆₈₁	96.6	88.8	13 050	46	28 100	100	7.2
V13An	81.2	73.2	-4000	14	-25 000	89	4.1
D-V13A	81.2	73.2	4000	14	25 050	89	4.1
V13K	74.9	64.7	-1400	5	-27 000	96	2.1
$D-V13K_{D}$	74.9	64.7	1500	5	26 950	96	2.1

Table 2: Biophysical data of peptide analogs

^aPeptide sequences are shown in Table 1.

^bPeptides are ordered by decreasing retention time ($t_{\rm R}$) in RP-HPLC at pH 2 at temperatures of 5 and 80 °C which is a measure of overall hydrophobicity.

^cThe mean residue molar ellipticities [θ]₂₂₂ (deg/cm²/dmol) at wavelength 222 nm were measured at 5 °C in benign conditions (100 mm KCl, 50 mm PO₄, pH 7.4) or in benign buffer containing 50% TFE by circular dichroism spectroscopy. The negative values in molar ellipticity denote the left-handed helices and the positive values denote the right-handed helices.

 d The helical content (in percentage) of a peptide relative to the molar ellipticity value of the peptide D-V₆₈₁ in the presence of 50% TFE.

 ${}^{\circ}P_{A}$ denotes the dimerization parameter of each peptide during the RP-HPLC temperature profiling, which is the maximal retention time difference of [$(t_{R}^{t} - t_{R}^{5}$ for peptide analogs) – $(t_{R}^{t} - t_{R}^{5}$ for control peptide C)] within the temperature range, and $(t_{R}^{t} - t_{R}^{5})$ is the retention time difference of a peptide at a specific temperature (*t*) compared with that at 5 °C.

TFE, trifluoroethanol; RP-HPLC, reversed-phase high-performance liquid chromatography.

on peptide retention behavior, i.e. a linear decrease in peptide retention time with increasing temperature because of greater solute diffusivity and enhanced mass transfer between the stationary and mobile phases at higher temperatures (34). Thus, after normalization to the retention times of peptide C, the retention behavior of the peptides represents only peptide self-association ability. Note that the higher the P_A value, the greater the self-association ability. The order of peptide self-association ability of the three pairs of peptide enantiomers is identical to the order of peptide hydrophobicity, i.e. V₆₈₁/D-V₆₈₁ have the highest oligomerization ability (e.g. dimerization) in solution among the three pairs of peptide enantiomers ($P_A = 7.2$, Table 2); in contrast, V13A_D/D-V13A_L showed a weaker ability to self-associate when compared with $V_{681}/D-V_{681}$ $(P_{\rm A} = 4.1, \text{ Table 2}); V13K_{\rm L}/\text{D-V13K}_{\rm D}$ exhibited the lowest self-association ($P_A = 2.1$, Table 2). As shown in Table 2, it is also clear that the peptide retention times at 80 °C are dramatically lower than those at 5 °C. Apart from the decrease in retention time because of the general temperature effects noted above, unraveling of the α -helix will also occur with increasing temperature, resulting in the loss of the non-polar face of the amphipathic α -helical peptides and, hence, reduced retention times as the peptides become increasingly random coils.

Hemolytic activity

The hemolytic activity of the peptides against human erythrocytes was determined as the maximal peptide concentration that produces no hemolysis after 18 h of incubation at 37 °C. The parent peptide, V₆₈₁, exhibited the strongest hemolytic activity with a value of 7.8 μ g/mL, compared with peptides V13K_L and V13A_D (250.0 μ g/mL and 31.3 μ g/mL, respectively; Table 3). The activity of the D-enantiomers was quantitatively equivalent to that of L-enantiomers.

In our previous study (20), the hemolytic activities of peptides V_{681} , V13A_D and V13K_I were determined as 15.6 μ g/mL, 250.0 μ g/mL and >250.0 µg/mL, respectively, following incubation for 12 h at 37 °C instead of the 18 h incubation of the present study. Hence, in order to explore the relationship between hemolysis and incubation time, a hemolysis time study was carried out to investigate the extent of hemolysis during different periods of incubation and different peptide concentrations. As illustrated in Figure 4, a hemolysis time study was carried out over 8 h at peptide concentrations of 8, 16, 32, 64, 125, 250 and 500 μ g/mL. The percentage hemolysis of cells was determined spectrophotometrically by comparison with the complete hemolysis of cells in water. As illustrated in Figure 4, peptides V₆₈₁, D-V₆₈₁, V13A_D, and D-V13A_L exhibited an increase in erythrocyte hemolysis with increasing incubation time at most peptide concentrations. Significantly, peptides $V13K_L$ and D-V13K_D showed negligible (<10%) or no hemolysis against human red blood cells after 8 h even at the extremely high peptide concentration of 500 μ g/mL. As illustrated in Figure 4, it is clear that the enantiomeric peptide pair V681/D-V681 exhibit cell lysis to a significantly greater extent than the V13A_D/D-V13A_L peptide pair. Thus, 100% hemolysis was already apparent after just 1 h incubation for V₆₈₁ and D-V₆₈₁ at a peptide concentration of 500 μ g/mL, whilst this value was approximately 40% for peptides V13A_D and D-V13A_I. In addition, while a small increase in hemolysis was apparent for V_{681} and D-V₆₈₁ at a peptide concentration of just 8 μ g/mL, negligible hemolysis was seen for $V13A_D$ and $D-V13A_L$ at a peptide concentration of 32 μ g/mL. It is interesting to note that for our lead compounds, peptides $V13A_D/D-V13A_L$ and $V13K_L/D-V13K_D$, D-enantiomers generally exhibited slightly weaker hemolytic activity compared with L-enantiomers at most peptide concentrations, the only exception being V13A_D and D-V13A_I at 500 μ g/mL, where the activity of the L-peptide is greater at <4 h incubation but weaker from 4 h onwards.



Figure 3: Peptide retention behavior during reversedphase high-performance liquid chromatography (RP-HPLC) with increasing temperature. Column and conditions: RP-HPLC, SB-C₈ column (150 \times 2.1 mm l.D.; 5 μ m particle size, 300 Å pore size), linear A-B gradient (0.5% acetonitrile/min) at a flow rate of 0.35 mL/min, where eluent A is 0.2% aqueous trifluoroacetic acid (TFA) and eluent B is 0.2% TFA in acetonitrile. The retention times of the peptides during the change of temperature are shown in panel (A). In panel (B), the retention time of peptides are normalized to 5 °C through the expression $(t_{\rm B}^t - t_{\rm B}^5)$, where $t_{\rm B}^t$ is the retention time at a specific temperature of an antimicrobial peptide or the random coil peptide, and $t_{\rm B}^5$ is the retention time at 5 °C. In panel (C), the retention behavior of the peptides was normalized to that of the random coil peptide C through the expression $(t_{\rm R}^t - t_{\rm R}^5$ for peptides) minus $(t_{\rm R}^t - t_{\rm R}^5$ for C). In panels (A-C), the symbols used are: circles for V_{681} and D- V_{681} , diamonds for V13AD and D-V13AL, squares for V13K_L, D-V13K_D and triangles for random coil peptide C.

Peptide antimicrobial activity and therapeutic index against Pseudomonas aeruginosa strains

Pseudomonas aeruginosa strains used in this study are a diverse group of clinical isolates from different places in the world. Antibiotic susceptibility tests show that these *P. aeruginosa* strains share similar susceptibility to most antibiotics except that there is about a 64-fold difference for the range of ciprofloxacin susceptibility (Table 3). Antimicrobial activities of peptide enantiomers against *P. aeruginosa* strains are shown in Table 3. In general, the antimi-

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crobial activity of L- and D-enantiomers against *P. aeruginosa* varied within fourfold. CP204, a *P. aeruginosa* strain from cystic fibrosis patients, is apparently the most susceptible strain to these peptides with MIC values from 7.8 to 31.3 μ g/mL. The geometric mean MIC values from six *P. aeruginosa* strains were calculated to provide an overall evaluation of antimicrobial activity of these peptides against *P. aeruginosa*. Fold improvement was calculated by comparing the geometric mean MIC value of each peptide to that of the parent peptide V₆₈₁. It is clear that, in most cases, all-D-peptides exhibited slightly better antimicrobial activity than their L-enantiomers.

The therapeutic index is a widely accepted parameter to represent the specificity of antimicrobial reagents. It is calculated by the ratio of MHC (hemolytic activity) and MIC (antimicrobial activity); thus, larger values of therapeutic index indicate greater antimicrobial specificity. The therapeutic indices of the peptides against *P. aerugi*nosa are shown in Table 3. By replacing L-valine with D-alanine or L-lysine, we significantly increased the therapeutic index against P. aeruginosa strains by 4.7- (peptide V13A_D) and 16.7-fold (peptide V13K₁). In addition, by making enantiomeric D-peptides, we further improved the therapeutic index against P. aeruginosa by 7.3- (peptide $D-V13A_1$) and 26.7-fold (peptide $D-V13K_D$) as measured by the standard microtiter dilution method. Peptide D-V13KD showed no hemolysis at 500 μ g/mL after 8 h in our time study (Figure 4) which is a more stringent test of hemolytic activity. The hemolytic activities of V₆₈₁ were not different when determined by the standard microtiter dilution method at 18 h or by the time study at 8 h using a peptide concentration of 500 μ g/mL (Table 3 and Figure 4). Thus, using a peptide concentration of 500 μ g/mL, the MHC value of $D-V13K_D$ increased the fold improvement of the therapeutic index compared with V_{681} to 53-fold. Our overall conclusion is that the key differences in biologic activities lie in the much more dramatic variations in hemolytic activity (MHC values) because the differences in antimicrobial activity between the six peptides are relatively minor (Table 3). Thus, the dramatic increases in therapeutic indices arise from the difference in hemolytic activity.

Peptide antimicrobial activity and therapeutic index against various gram-negative bacteria

Table 4 shows peptide antimicrobial activity and therapeutic indices against six different gram-negative bacterium strains. It is clear to see that, in general, all-L-peptides have equivalent antimicrobial activity against gram-negative bacteria compared with their D-enantiomers, because MIC values of L- and D-enantiomers are all within a twofold difference. In Table 4, geometric mean MIC was calculated to provide an overall view of antimicrobial activity of the peptides against gram-negative bacteria and was also used to calculate the therapeutic index. Peptides $V13K_1$ and $D-V13K_D$ both show a 40-fold improvement in therapeutic index against gram-negative bacteria compared with peptide V₆₈₁ as measured by the standard microtiter dilution method. For this improvement in the therapeutic index of peptides against gram-negative bacteria, peptide hemolytic activity is again the key factor. Using our most stringent test of hemolytic activity, the MHC value of 500 μ g/mL for peptide D-V13KD showed no hemolysis after 8 h in our time study (Figure 4) and the fold improvement of the therapeutic index compared with V681 was 80-fold.

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Table 3: Hemolytic activities of peptides and MIC against Pseudomonas aeruginosa strains

	Hemolytic activit	у	Antimicr	obial activit	y						Therapeutic i	ndex
			MIC (µg,	/mL) ^c								
Peptide	MHC $(\mu g/mL)^a$	Fold ^b	PAO 1	WR 5	PAK	PA 14	M 2	CP 204	GM^d	Fold ^e	MHC/MIC ^f	Fold ^g
V ₆₈₁	7.8	1	15.6	62.5	31.3	62.5	15.6	15.6	27.8	1.0	0.3	1.0
D-V ₆₈₁	7.8	1	15.6	31.3	31.3	15.6	31.3	31.3	24.8	1.1	0.3	1.0
V13An	31.3	4	15.6	62.5	62.5	31.3	7.8	7.8	22.1	1.3	1.4	4.7
D-V13A	31.3	4	7.8	31.3	31.3	7.8	15.6	7.8	13.9	2.0	2.2	7.3
V13K	250.0	32	31.3	250.0	125.0	62.5	31.3	7.8	49.6	0.6	5.0	16.7
D-V13K₀ Ciprofloxacin ⁱ	250.0 ^h	32	15.6 4	62.5 1	125.0 4	15.6 8	31.3 8	15.6 64	31.2	0.9	8.0	26.7 (53.4) ^h

^aMHC denotes the maximal peptide concentration that produces no hemolysis after 18 h in the standard microtiter dilution method.

^bDenotes the fold decrease in hemolytic activity compared with the parent peptide V₆₈₁.

^cMIC is minimal inhibitory concentration that inhibited growth of six *P. aeruginosa* strains. MIC is given based on three sets or more of determinations.

^dDenotes the geometric mean of a diverse group of *P. aeruginosa* clinical isolates.

^eDenotes the fold improvement in antimicrobial activity (geometric mean data) compared with the parent peptide V₆₈₁.

^fTherapeutic index is the ratio of the MHC value (μ g/mL) over the geometric mean MIC value (μ g/mL). Large values indicate greater antimicrobial specificity. ^gDenotes the fold improvement in therapeutic index compared with the parent peptide V₆₈₁.

^hIn the hemolysis time study, D-V13K₀ showed no hemolysis at 500 μ g/mL for 8 h, which would increase the fold improvement to 53.4-fold in the therapeutic index.

ⁱRelative susceptibility of these *Pseudomonas* strains to ciprofloxacin with the number 1 denoting the most susceptible strain.



Figure 4: Time study of peptide hemolysis of human red blood cells. Closed symbols were used for L-peptides and open symbols were used for D-peptides. Different symbols were used to represent the different concentrations of peptides during the hemolysis study.

Peptide antimicrobial activity and therapeutic index against various gram-positive bacteria and a fungus

The antimicrobial activity against gram-positive bacteria and a fungus of L- and D-enantiomeric peptides is compared in Table 5. It is clear that our peptides were effective in killing the gram-positive bacteria tested. The least sensitive gram-positive bacterium to the enantiomeric peptides was *Enterococcus faecalis*. The effectiveness of the peptides to lyse cells of the fungus *Candida albicans* was similar to *E. faecalis*. For gram-positive bacteria and the fungus, D-peptides again generally showed the same or better antimicrobial activity than L-enantiomers. Although the geometric mean MIC values of peptides V13K_L/D-V13K_D were among the two lowest, because of the high MHC values (poor or no hemolytic activity), the therapeutic indices of V13K_L and D-V13K_D were the highest against gram-positive bacteria and the fungus. Similar to that seen against gram-negative bacteria, the therapeutic index of peptide D-V13K_D was increased 34.7- and 16.7-fold against gram-positive bacteria and *C. albicans*, respectively, because of the poor or no hemolytic activity as measured by the standard microtiter dilution method. Using our most stringent test of hemolytic activity, the MHC value of 500 μ g/mL for peptide D-V13K_D showed no hemolysis after 8 h in our time study (Figure 4) and the fold improvement of the therapeutic indices compared with V₆₈₁ were 69- and 33-fold against gram-negative bacteria and fungus, respectively.

Proteolytic degradation study

Figure 5 illustrates the stability of L- and D-enantiomeric peptides in the presence of trypsin at a molar ratio of 20 000:1 or 20 mM: 1 μ M (peptide:trypsin) at 37 °C. The assay was carried out for 8 h; however, only the stability of the peptides in the first 60 min is illustrated in Figure 5, due to the reason that all three L-peptides were 100% digested by trypsin after 1 h of incubation at 37 °C. It is interesting to see that peptide V₆₈₁ showed slightly more resistance against trypsin than V13K_L and V13A_D; in contrast, V13K_L and

Table 4: Antimicrobial (MIC) and hemolytic (MHC) activities of peptide analogs against Gram-negative bacteria and human red blood cells

	MHC $^{\rm a}$ (μ g/mL)	$MIC^{\mathtt{b}}$ (μ g/mL	_)							Therapeutic	index ^f
Peptides	hRBC	<i>E. coli</i> C498 (wt) ^c	<i>E. coli</i> C500 (abs) ^c	<i>St</i> C587 (wt) ^c	<i>St</i> C610 (abs) ^c	<i>Pa</i> H187 (wt) ^c	<i>Pa</i> H188 (abs) ^c	GM ^d	Fold ^e	MHC/MIC	Fold ^g
V ₆₈₁	7.8	6.3	3.1	12.5	3.1	6.3	3.1	5.0	1.0	1.6	1.0
D-V ₆₈₁	7.8	6.3	3.1	12.5	6.3	6.3	6.3	6.3	0.8	1.2	0.8
V13AD	31.3	3.1	1.6	12.5	3.1	3.1	1.6	3.1	1.6	10.1	6.3
D-V13A	31.3	3.1	3.1	12.5	3.1	3.1	3.1	3.9	1.3	8.0	5.0
V13K	250.0	3.1	3.1	6.3	3.1	6.3	3.1	3.9	1.3	64.1	40.1
D-V13K _D	250.0 ^h	3.1	3.1	6.3	3.1	6.3	3.1	3.9	1.3	64.1	40.1 (80.2) ^h

^aHemolytic activity (maximal peptide concentration with no hemolysis after 18 h) was determined on human red blood cells (hRBC).

^bAntimicrobial activity (minimal inhibitory concentration) is from three sets of determinations.

^cE. coli, Escherichia coli, St, Salmonella typhimurium; Pa, Pseudomonas aeruginosa; wt, wild-type laboratory strain; abs, antibiotic-sensitive strain.

^dGM denotes the geometric mean of MIC values from all six microbial strains in this table.

^eDenotes fold improvement in the geometric mean of MIC compared with parent peptide V_{681} .

^fTherapeutic index = MHC (μ g/mL)/geometric mean of MIC (μ g/mL). Larger values indicate greater antimicrobial specificity.

 g Denotes fold improvement in the therapeutic index compared with parent peptide V₆₈₁.

^hIn the hemolysis time study, p-V13K_p showed no hemolysis at 500 μ g/mL for 8 h which would increase the fold improvement in the therapeutic index to 80.2-fold.

V13A_D generally showed a similar trend of degradation in the presence of trypsin. All three peptides were 86–97% digested after 30 min. In sharp contrast, D-peptides were completely stable (no degradation) to trypsin even at a 10-fold higher molar concentration ratio of trypsin versus peptide (2 mM peptide:1 μ M trypsin) for 8 h (data not shown).

Discussion

Enantiomers are non-superimposable mirror images of one another. When a peptide is composed of all-D-amino acids, it becomes the Denantiomer of the all-L-peptide in the same sequence. In such cases, the original left-handed helix of the L-peptide will be present as a right-handed helix in its all-D-enantiomeric form, as demonstrated by the CD data illustrated in Figure 2. The L- and D-enantiomeric pairs of V₆₈₁ exhibited <50% α -helical structure in benign buffer while the two analog pairs exhibited little or negligible α -helical structure in benign buffer; however, all three enantiomeric pairs were fully helical in the presence of 50% TFE. This characteristic ensures α -helical structure when the peptides interact with the biomembrane, as TFE has been frequently employed as a mimic of the hydrophobic environment characteristic of biomembranes (18,20). We believe having no α -helical structure in benign medium and inducible α -helical structure in the hydrophobic environment of the membrane is a critical aspect for antimicrobial activity.

In the present study, we have clearly shown that each enantiomeric peptide pair has the same activities against prokaryotic and eukaryotic cell membranes which suggest that the sole target for these antimicrobial peptides is the cell membrane. We previously proposed a model (20) to explain the peptide specificity between prokaryotic and eukaryotic membranes due to the different compositions of lipids between prokaryotic and eukaryotic membranes. Eukaryotic cell membranes, in contrast to prokaryotic membranes, are generally characterized by zwitterionic phospholipids, a relatively large amount of cholesterol and sphigomyelin, and the absence of the high, inside-negative transmembrane potential of prokaryotic membranes (35-38). This model proposes that hemolysis of eukaryotic cells requires the peptides to be inserted into the hydrophobic core of the membrane, perpendicular to the membrane surface, and interaction of the non-polar face of the amphipathic α -helix with the hydrophobic lipid core of the bilayer. The peptide may thus form transmembrane channels/pores and the hydrophilic surfaces point inward, producing an aqueous pore ('barrel-stave' mechanism) (39). In contrast, antimicrobial activity in prokaryotic cells, while maintaining specificity, requires the peptide to lie at the membrane interface parallel with the membrane surface and interaction of the non-polar face of the amphipathic α -helix with the hydrophobic component of the lipid and interaction of the positively charged residues with the negatively charged head groups of the phospholipid ('carpet' mechanism) (40,41). What dictates the two different modes of interaction is the difference in lipid composition of prokaryotic and eukaryotic membranes. We refer to this mode of interaction of antimicrobial peptides which combines the above two mechanisms as a 'membrane discrimination mechanism'.

Using this model, it is easy to understand why peptide V13K₁ and $D-V13K_D$ of the present study are non-hemolytic but at the same time possess excellent antimicrobial activity compared with the native sequence V_{681} (V₁13) or D-V₆₈₁ (V_D13). Thus, the single substitution of Lys for Val at position 13 (V13K₁ and D-V13K_D) in the center of the non-polar face disrupts the hydrophobic surface due to the presence of the positive charge, preventing the peptide from penetrating the bilaver as a transmembrane helix in eukarvotic cells. The peptide is then excluded from the bilayer and, hence, is non-hemolytic. In prokaryotic cells, the peptide is also excluded from penetrating the bilayer as a transmembrane helix but this is not required for excellent antimicrobial activity. Instead, the peptide can enter the interface region of the bilayer where disruption of the peptide hydrophobic surface by Lys can be tolerated and antimicrobial activity maintained. These are consistent with the results of previous studies of model membranes which demonstrated that pore formation mechanism ('barrel-stave' mechanism) was used by

	MHC ^a (µg/mL)	MIC ^b (µg/m	IL)							Therapeutic	index ^e	MIC ^b (µg/mL)	Therapeutic	index ^f
Peptides	hRBC	<i>S. aureus</i> 622 (wt) ^c	<i>S. aureus</i> 623 (methR) ^c	S. epidermidis C621 (wt) ^c	<i>B. subtilis</i> C971 (wt) ^c	<i>E. faecalis</i> C625 (wt) ^c	<i>C. xerosis</i> C875 (wt) ^c	GM ^d	Fold ^d	MHC/MIC	Fold ^g	C. albicans	MHC/MIC	Fold ⁹
V ₆₈₁	7.8	3.1	6.3	6.3	3.1	6.3	3.1	4.4	1.0	1.8	1.0	12.5	0.6	1.0
D-V ₆₈₁	7.8	6.3	6.3	1.6	1.6	6.3	3.1	3.5	1.3	2.2	1.2	6.3	1.2	2.0
V13A _D	31.3	1.6	3.1	1.6	3.1	12.5	1.6	2.8	1.6	11.2	6.2	12.5	2.5	4.2
D-V13A	31.3	3.1	3.1	1.6	1.6	12.5	1.6	2.8	1.6	11.2	6.2	12.5	2.5	4.2
V13K	250.0	12.5	12.5	6.3	3.1	50.0	1.6	7.9	0.6	31.6	17.6	50.0	5.0	8.3
D-V13K _D	250.0 ^h	6.3	6.3	3.1	1.6	12.5	1.6	4.0	1.1	62.5	34.7 (69.4) ^h	25.0	10.0	16.7 (33.4

Antimicrobial and antifungal activities (minimal inhibitory concentration) are from three sets of determinations.

aureus, Staphylococcus aureus, S. epidermidis, Staphylococcus epidermidis, B. subtilis, Bacillus subtilis, E. faecalis, Enterococcus faecalis, C. xerosis, Corynebacterium xerosis, C. albicans, Candida albicans, xtu, S.

⁴3M denotes the geometric mean of MIC values from all six gram-positive microbial strains in this table. Fold denotes fold improvement in the geometric mean MIC compared with parent peptide V₆₈₁ wild-type strain; methR, methicillin-resistant clinical strain.

^Therapeutic index = MHC (μ g/mL/geometric mean MIC (μ g/mL). Larger values indicate greater antimicrobial specificity

Denotes the therapeutic index of the peptides against a fungus C. albicans.

³Denotes fold improvement in the therapeutic index compared with parent peptide V₆₈₁.

¹In the hemolysis time study. D-V13K₀ showed no hemolysis at 500 µg/mL for 8 h which would increase the fold improvement in the therapeutic index to 69.4- and 33.4-fold for gram-positive bacteria and C. albicans, respectively.



Figure 5: Peptide stability to proteolysis by trypsin. Closed symbols were used for L-peptides and open symbols were used for D-peptides. Circles denote V_{681} and D- $V_{681},$ squares denote $V13K_L$ and D-V13K_D, and diamonds denote V13A_D and D-V13A_L.

antimicrobial peptides in zwitterionic membranes whilst a detergent-like mechanism ('carpet' mechanism) for the peptides was shown when the peptides interact with negatively charged membranes (42,43). Thus, the position of the Lys residue in the center of the non-polar face becomes a critical factor in maintaining specificity between prokaryotic and eukaryotic membranes. This approach was previously used by our laboratory with cyclic β -sheet peptides (44-47).

The technique of RP-HPLC temperature profiling to monitor molecule self-association has been applied to several types of polypeptides, including cyclic β -sheet peptides (29), monomeric α -helices and α -helices that dimerize (20,28), including α -helices that dimerize to form coiled-coils (33). We believe that peptide self-association (i.e. the ability to dimerize) in aqueous solution is a very important parameter to understand the mechanism of action of antimicrobial peptides. *α*-Helical antimicrobial peptides are amphipathic; thus, if the self-association ability of a peptide (forming dimers by interaction of the two non-polar faces of two molecules) is too strong in aqueous media, it could decrease the ability of the peptide monomers to dissociate, pass through the cell wall of micro-organisms and penetrate into the biomembranes to kill target cells. In the present study, there is a direct correlation of the ability of peptides to dimerize and specificity, i.e. disruption of dimerization generates specificity between eukaryotic and prokaryotic cells. As shown in Table 2, the P_A values of peptides derived from their temperature profiling data (Figure 3) reflect the ability of the amphipathic α -helices to associate/dimerize. Clearly, V_{681} and D- V_{681} , because of their uniform non-polar faces, show the greatest ability to associate in aqueous solution and lowest specificity or the strongest ability to lyse human erythrocytes. This result supports the view that a peptide with a fully accessible non-polar face tends to form pores/ channels in the membranes of eukaryotic cells. In the case of V13A_D and D-V13A_L, the introduction of D-Ala and L-Ala into

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Table 5: Antimicrobial (MIC) and hemolytic (MHC) activities of peptide analogs against Gram-positive bacteria, fungus and human red blood cells

all-L- and all-D-amino acid peptides, respectively, disrupts α -helical structure and, thus, lowers dimerization ability relative to V_{681} and D-V₆₈₁ and improves specificity. The introduction of Lys into nonpolar position 13 of $V13K_L$ and D-V13K_D lowers this association ability even further and further improves specificity. Thus, the lack of ability of a peptide to oligomerize or dimerize, as exemplified by its $P_{\rm A}$ value, is an excellent measure of the peptide's ability to be nonhemolytic concomitant with maintenance of sufficient hydrophobicity of the non-polar face to ensure antimicrobial activity. It is important to note that, in the present study, D-peptides exhibited the same self-association ability as their L-enantiomers: thus, similar biologic activities can be expected. This is shown by the fact that hemolytic activity and antimicrobial activity of D-peptides against human red blood cells and microbial cells, respectively, were quantitatively equivalent to that of the L-enantiomers, further demonstrating that there is no chiral selectivity by the membrane or other stereoselective interactions in the cytoplasm during hemolytic and antimicrobial activities.

Because of the dramatic different results on hemolytic activity of peptide V13A_D in the previous (20) and the present study (250 μ g/ mL after 12 h versus 31.3 μ g/mL after 18 h, respectively), using the standard microtiter dilution method (see Materials and Methods), it became apparent that an investigation of the relationship between hemolysis and time was required. It is noteworthy that there is no universal protocol of determination of hemolytic activity, which makes it difficult to compare data from different sources. For example, some researchers use 4 h of incubation and take the minimal concentration of peptide to give 100% hemolysis as peptide hemolytic activity (21,48); in contrast, some use 12 h or longer (e.g. 18 h in this study) of incubation and take the maximal concentration of peptide to give no hemolysis as peptide hemolytic activity [(12,20) and the present study]. Hence, the hemolysis time study is important to understand the process of erythrocyte lysis. It is clear that the degree of cell lysis is correlated with time, which may be the main reason for the different values of hemolytic activity of V13AD in the two studies. Hence, we have established a stringent criterion for toxicity, which is no hemolysis at a peptide concentration of 500 μ g/mL after 8 h. We believe that this time study at this very high peptide concentration gives a much more accurate evaluation of hemolytic activity and this method should be established as the gold standard test.

As mentioned before, *P. aeruginosa* is a family of notorious gramnegative bacterial strains which are resistant to most current antibiotics; thus, it is one of the most severe threats to human health (23–25). Only a few antibiotics are effective against *Pseudomonas*, including fluoroquinolones (49), gentamicin (50), and imipenem (51), and even these antibiotics are not effective against all strains. In this study, it is encouraging to see that our lead peptides V13A_D and V13K_L are effective against a diverse group of *P. aeruginosa* clinical isolates. Peptide D-V13A_L exhibited the highest antimicrobial activity against *P. aeruginosa* strains; in contrast, D-V13K_D has the best overall therapeutic index due to its lack of hemolytic activity. It has to be pointed out here that, in this study, MIC values for *P. aeruginosa* and other gram-negative and gram-positive bacteria were determined in two different collaborating laboratories; in addition to different media used, the inocu-

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lum numbers of cells were also different (see details in the Materials and Methods), which may cause some variations of MIC values of *P. aeruginosa* strains when comparing the results reported in Tables 3 and 4.

It is important to note that, in general, there is no significant difference in peptide antimicrobial activities against P. aeruginosa strains, other gram-negative and gram-positive bacteria and a fungus between L- and D-enantiomeric peptides, or among peptides with different amino acid substitutions, i.e. V_{681} , V13A_D, and V13K₁ (Tables 3–5). This observation provides understanding of the mechanism of action of *a*-helical antimicrobial enantiomeric peptides as follows: there is a dramatic difference in peptide hydrophobicity at position 13 between Val and Lvs. The Lvs disrupts the continuous non-polar surface because of the positive charge and causes the peptide to locate in the interface region of the microbial membrane. This supports the view that the 'carpet' mechanism is essential for strong antimicrobial activity, i.e. for both L- and D-peptide enantiomers, the peptides kill bacteria by a detergent-like mechanism, without penetrating deeply into the hydrophobic core of membrane.

Based on the peptide degradation study, all-D-peptides were totally resistant to enzymatic digestion; hence, this may explain the slightly higher antimicrobial activity of D-peptides than that of their L-enantiomers against *P. aeruginosa* and gram-positive bacteria. The relatively high susceptibility of L-peptides to trypsin is no doubt due to the presence of multiple lysine residues in sequences, i.e. six lysines for V₆₈₁ and V13A_D, seven lysines for V13K_L, resulting in the fast degradation of the L-peptides in 30 min even at a molar ratio of 20 000:1 (peptide:trypsin).

In conclusion, by comparing the biophysical and biologic properties of L- and D-enantiomeric peptides, we report here that L- and D-enantiomeric peptide pairs behaved the same in self-association ability in solution, had the same hemolytic activity against human red blood cells, and exhibited similar antimicrobial activity against P. aeruginosa strains, and other gram-negative and gram-positive bacteria and a fungus. No chiral selectivity was found in the antimicrobial and hemolytic activities of the peptides. In this study, our membrane discrimination mechanism has proved to be a good explanation of the mechanism of action for both L- and D-enantiomeric peptides. It is important to note that peptide $D-V13K_D$ has demonstrated dramatic improvements in therapeutic indices compared with the parent peptide V₆₈₁ by 53-fold against P. aeruginosa strains, 80-fold against gram-negative bacteria, 69-fold against gram-positive bacteria, and 33-fold against C. albicans. The proteolytic stability of D-V13KD, its broad spectrum activity and lack of hemolytic activity demonstrate its clinical potential as a new therapeutic.

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