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Identification and bioactivity evaluation of two novel temporins from the skin secretion of the European edible frog, Pelophylax kl.esculentus

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Abstract

Two novel members of the temporin family, named temporin-1Ee (FLPVIAGVLSKLFamide) and temporin-1Re (FLPGLLAGLLamide), whose biosynthetic precursor structures were deduced from clones obtained from skin secretion-derived cDNA libraries of the European edible frog, Pelophylax kl. esculentus, were obtained by ‘shotgun’ cloning. Deduction of the molecular masses of each mature processed peptide from respective cloned cDNAs was used to locate respective molecules in reverse-phase HPLC fractions of secretion. Temporin-1Ee (MIC=10µM) and temporin-1Re (MIC=60µM) were both found to be active against Gram-positive Staphylococcus aureus, but retaining a weak haemolytic activity. Single-site substitutions can dramatically change the spectrum of activity of a given temporin. Compared with temporine-1Ec, just one chemically-conservative substitution (Val8 instead of Leu8), temporin-1Ee bearing a net charge of +2 displays broad-spectrum activity with particularly high potency on the clinically relevant Gram-negative strains, Escherichia coli (MIC=40µM). These factors bode well for translating temporins to be potential anti-infective drug candidates.
Keywords: Amphibian; Antimicrobial Peptide; Temporins; Cloning

Introduction

Temporins, containing 10-13 amino acid residues, are considered to be among the smallest AMPs found in Nature. They were initially described as the Vespa-like peptides, due to their sequence similarities with chemotactic and histamine-releasing peptides isolated from the venom of wasps of the genus Vespa, and were first identified in the skins of the European edible frog, *Pelophylax kl. esculentus* and the Asian frog, *Rana erythraea*. Afterwards, a family of 10, structurally-related peptides with antibacterial and antifungal properties were isolated from the electrically-stimulated skin secretions of the European common red frog, *Rana temporaria*, and called temporins A to L, in reflection of their source (1). Compared with other ranid frog skin AMPs that have a C-terminal heptapeptide ring constructed by a disulphide bridge, the temporins are C-terminally α-amidated and stabilised by helix-forming residues such as leucine, alanine and lysine. Due to their basic residues, temporins usually carry a net charge ranging from 0 to +4 at neutral pH and have a potent activity against Gram-positive bacteria including methicillin-resistant *S. aureus* (MRSA) with little toxicity to mammalian cells (2,3). However, since temporin L is the first member of this family isolated which exhibit broad spectra of activity against both Gram-positive and Gram-negative bacteria (Giacometti et al 2006), a few new members of the temporin family, such as temporin-Sha, temporin-T1 and temporin-1DRa, have been found to display similar antimicrobial activities to temporine L(4, 5).

Naturally occurring AMPs differ dramatically in various interrelated peptide parameters, including size, net charge, hydrophobicity, amphipathicity, degree of structure formation and flexibility (6-8), thus the selectivity and mechanism of antimicrobial activities remain unknown. It is well accepted that differences in lipid composition of membranes among microorganisms play key roles in the selectivity of AMPs toward microbial cells (2), and
although the mechanism of action of temporins have been only scarcely evaluated, most are supposed to have a classical membrane perturbing activity through a barrel-stave mechanism of action. In contrast, temporin L is presumed to insert into bacterial cell membranes through the formation of pore-like openings and interact with essential anionic targets inside the cells (1-3, 9-10). Temporins as short chain peptides of simple structure are considered to be good templates for new drug design.

Herein, two novel antimicrobial peptides have been identified in the skin secretion of the European Edible frog, *Pelophylax kl. esculentus*, through initial molecular cloning of skin secretion-derived cDNAs. Deduced peptides were subsequently identified in fractions of skin secretion, chemically-synthesised and subjected to functional analyses. Sequence characterisation of the mature peptides demonstrated that they belonged to the temporin family found predominantly in the skin secretions of ranid frogs. Thus, these two peptides were named temporin-1Ee and temporin-1Re. Temporin-1Ee, which is an analogue of temporin-1Ec, has broad-spectrum antimicrobial activity at low micromolar concentrations. Temporin-1Re, with a net charge of +1, is weakly active against the Gram-positive bacterium, *S. aureus*, but is devoid of activity against Gram-negative bacteria and yeast.
Materials and methods

Preparation of *P. kl.esculentus* skin secretion

*P. kl.esculentus* (*n* = 30) were obtained from a commercial source (Le QuerruyCellier, Notre Dame de Monts, France) and had been captive bred. The frogs were adults on receipt and were settled into their new surroundings for 3 months prior to secretion harvesting. They were maintained in our purpose-designed amphibian facility at 15–20 °C under a 12 h/12 h light/dark cycle and fed multivitamin-loaded crickets three times per week. Skin secretion was obtained from dorsal skin by mild transdermal electrical stimulation (5V; 3 ms pulses) for 30 s. The skin secretion was washed from the skin using deionized water, snap-frozen in liquid nitrogen, and lyophilized. Lyophilizate was stored at −20 °C prior to analysis.

“Shotgun” cloning of *Pelophylax kl. esculentus* skin secretion-derived cDNA

Five milligrams of lyophilised skin secretion were dissolved in 1ml lysis/binding buffer (Dynal, UK) to stabilise mRNA. Polyadenylated mRNA was trapped and isolated by means of magnetic oligo-dTDynabeads as described by the manufacturer (Dynal, UK). The trapped template polyadenylated mRNA was employed to manufacture a cDNA library using reverse transcriptase and a sample of this library was subjected to 5′- and 3′-rapid amplification of cDNA ends (RACE) procedures to obtain full-length antimicrobial peptide precursor nucleic acid sequence data using a SMART-RACE kit (Clontech UK) following manufacturer's instructions. Briefly, the 5′-RACE reactions employed the forward primer (5′-ATGTTCACCATGAAGAAATC-3′) and the reverse primer (5′-CTATCCCACATCATAGAGACTTTCC-3′) were designed according to the nucleotide sequence of temporin precursor cDNAs obtained from other ranid frogs. The 5′-RACE reactions were purified and cloned using a pGEM-T vector system (Promega Corporation) and sequenced using an ABI 3730 automated sequencer (Applied Biosystems, Foster City, CA, USA).
Reverse phase HPLC fractionation of skin secretion

A further 1mg sample of lyophilised skin secretion was dissolved in 0.5ml of 0.05/99.5 (v/v) trifluoroacetic acid (TFA)/water and clarified of microparticulates by centrifugation. The supernatant was then subjected to HPLC using a gradient formed from 0.05/99.5 (v/v) TFA/water to 0.05/29.95/70.0 (v/v/v) TFA/water/acetonitrile in 240min at a flow rate of 1 ml/min. A Cecil Adept Binary HPLC system (Adept Technology, Inc. USA), fitted with an analytical column (a Jupiter 00G4052 semi-preparative C-5 column 25x1cm, Phenomenex, U.K.) was employed and this was interfaced with a Thermo Fisher Scientific LCQ Fleet electrospray ion-trap mass spectrometer. The effluent from the chromatographic column was flow-split with approximately 10% entering the mass spectrometer source and 90% directed towards a fraction collector. Dead volume between column and fraction collector was minimal (20μl). The molecular masses of polypeptides in each chromatographic fraction were further analysed using matrix-assisted laser desorption/ionisation, time-of-flight mass spectrometry (MALDI-TOF MS) on a linear time-of-flight Voyager DE mass spectrometer (Perseptive Biosystems, MA, USA) in positive detection mode using alpha-cyano-4-hydroxycinnamic acid as the matrix. Internal mass calibration of the instrument with known standards established the accuracy of mass determination as ±0.1%.

Peptide synthesis and purification

Following unequivocal establishment of the primary structure of natural temporin-1Ee and temporin-1Re, replicates were synthesised by solid-phase Fmoc chemistry using a PS3 automated peptide synthesiser (Protein Technologies, Tucson, AZ, USA). When the synthesis cycles were completed, the peptides were cleaved from the resins using 95/2.5/2.5 (v/v/v) TFA/TIPS/water for 6h, precipitated in ether over the next 24 h, washed exhaustively in six changes of ether and then allowed to completely dry over a further 24h. The peptides were then dissolved in a minimal quantity of 0.05/99.5, v/v, TFA/water, snap frozen in liquid
nitrogen and lyophilised. Degree of purity and authentication of structures of the synthetic peptides were determined using MALDI-TOF MS as previously described.

**Antimicrobial assays**

Antimicrobial activities of synthetic temporin-1Ee and temporin-1Re were assessed by determination of minimal inhibitory concentrations (MICs) using a standard Gram-positive bacterium *S. aureus*—NCTC 10788), a standard Gram-negative bacterium (*Escherichia coli*—NCTC 10418) and a standard pathogenic yeast (*Candida albicans* NCPF 1467). Temporin-1Ee and temporin-1Re were tested within the concentration range of 160–2.5μM and were initially dissolved as a stock solution of 200μM in sodium phosphate-buffered saline, pH 7.2, and subsequently diluted in Mueller–Hinton broth (MHB). Peptide concentrations in the range stated, were inoculated with microorganism cultures (105 colony forming units (CFU)/ml), and placed into 96-well microtiter cell culture plates. Plates were incubated for 18 h at 37°C in a humidified atmosphere. Following this, the growth of bacteria/yeast was determined by means of measuring optical density (OD) at λ=550nm by an ELISA plate reader (BioliseBioTek EL808). Minimal inhibitory concentrations (MICs) were defined as the lowest concentration at which no growth was detectable.
Results

Molecular cloning of the A biosynthetic precursor

Two different cDNAs, each encoding a novel peptide, named temporin-1Ee and temporin-1Re, respectively, were consistently and repeatedly cloned from the library during RACE-PCR experiments (Figure 1). Such experiments used R. esculenta skin mRNA as a template and specific oligonucleotides, designed to a conserved region (Signal peptide and 3′-UTR) of previously cloned preprotemporincDNAs from other ranid species. The cDNAs amplified were separated on agarose gels then isolated with the cloning vector which was transformed into host cells. The white and drug resistant bacterial colonies were chosen by random and subjected to nucleotide sequence analysis. Alignment of temporin-1Ee and -1Re nucleotide sequences and open-reading frame amino acid sequences, using the AlignX programme of the Vector NTI Bioinformatics suite (Informax), revealed a very high degree of primary structural similarity of both nucleic acid and amino acid sequence between the temporin-1Ee and -1Re (Figure 2). An NCBI BLAST search found that the novel temporin-1Ee showed at least 92% sequence identity with temporin-1Ec and peptide A1 and peptide B9 obtained from R. esculenta, previously (Table 2). In the precursor sequence, the first 22 amino acid residues encoded a putative signal peptide at the N-terminal followed by an acidic amino acid residue rich spacer peptide consisting of 21 amino acids for temporin-1Ee and 22 amino acids for temproin-1Re with a pair of classical basic residue -Lys-Arg- (-K-R-) propeptide convertase cleavage sites at its C-terminus. A single copy of a mature temporin sequence was found at the C-terminus just upstream of a Gly residue that serves as an amide donor for the C-terminal residue of the temporin. The nucleotide sequence of the temporin-1Ee and temporine-1Re precursors from the skin secretion of the European edible frog, Pelophylax kl.esculentus, have been deposited in the EMBL Nucleotide Sequence Database under the accession code KT437658 and KT437659.
Identification and structural analysis of mature temporin-1Ee and temporin-1Re in skin secretion fractions and subsequent chemical synthesis

The two novel temporins (temporin-1Ee: FLPVIAGVLSKLF-amide and temporin-1Re: FLPGLLAGLL-amide), deduced from cloned skin cDNAs, with computed molecular mass of 1402.08Da and 1012.32Da, respectively, were identified in HPLC fraction numbers 164 and 143, respectively (Figures 3 and 4). Both temporins were successfully obtained by the solid-phase chemical synthesis. The reverse phase HPLC chromatograms obtained from each synthesis following cleavage from the resin and deprotection, showed major peaks in each case representing approximately 95% of the synthetic mixture and these were subjected to MALDI–TOF mass analysis. In each case, molecular ions consistent with predicted molecular masses of respective novel temporins were observed indicating successful syntheses and confirming a high degree of purity (>95%) of the expected products.

Antimicrobial/hemolytic activity of temporin-1Ee and temporin-1Re

The abilities of temporin-1Ee and temporin-1Re peptides to inhibit the growth of the tested microorganisms employed are compared in Table 3. MICs obtained against S. aureus were 10 µM for temporin-1Ee and 60 µM for temporin-1Re. Moreover, temporin-1Ee displayed an inhibitory effect on the growth of the Gram-negative bacterium, E. coli, MIC was 40 µM, whereas temporin-1Re exhibit no effect. Both temporins were ineffective in inhibiting the growth of the yeast, C. albicans, at concentrations up to and including 160µM. Moreover, temporin-1Ee was found to possess no haemolytic activity at the MIC (10µM) determined for S. aureus. This experiment was repeated five times.
Discussion

The AMPs stored in the glands of frog skin display a complicated array of typical peptide physico-chemical attributes related to size, charge, hydrophobicity, degree of secondary structure formation and amphipathicity. Temporins have been intensely-studied due to their potent activity against Gram-positive bacteria, including clinical isolates of methicillin-resistant *Staphylococcus aureus* and vancomycin-resistant *Enterococcus faecium* and *Enterococcus faecalis* (9,12). However, due to their small size and low charge, the antimicrobial activity mechanisms of temporins are still unclear. The most recent research has provided some new insights to explain the mechanism of action of the temporins. Take temporin-L as an example. Temporin-L is a 13-mer peptide in which a tail-to-tail dimerisation can occur that then suits the barrel-stave model, binding to the vesicles surface, and undergoing a rapid local aggregation that causes a reorientation. Membrane insertion steps can only occur after this oligomerisation (12-14). We presume that temporin-1Ee, which has the same chain length as temporin-L and also possesses antimicrobial activity against both Gram-negative and Gram-positive bacteria, may act via a similar mechanism in which the carpet model could explain this temporin’s antimicrobial activity but the barrel-stave model could explain the haemolytic effect (1-3, 15).

Hence the variation in net charges (0 to +3), chain length (generally 10-14 residues), and amino acid sequences could influence the efficacy and selectivity of temporins against pathogens. Most members of the temporin family carry a low net charge of 0 or +1 at physiological pH and their chain lengths are restricted and small. Within the temporins, the higher the net positive charge (+2 or +3), the more potent they are in killing bacteria. Temporin-1Ee, with a net charge of +2, is more potent (MIC 10µM) than temporin-1Re (MIC 60µM), with a net charge of +1, in killing the model Gram-positive bacterium, *Staphylococcus aureus* (NCTC 10788). In contrast, due to most of the natural
temporinshaving a weak charge and small size, they generally have little or no effects on Gram-negative bacteria, fungi and yeasts (16-20). However, one of the prototype temporins, temporin L (13 residues, net charge +3), was found to exhibit broad-spectrum activity against many microorganisms, including Gram-positive bacteria, Gram-negative bacteria and yeast, even cancer cells and human erythrocytes. Many additional temporins subsequently discovered exhibit broad-spectrum antimicrobial activity, examples being temporin-1Sa (11) and temporin-1DRa (21-22). Temporin-1Ee, with a net charge of +2, has efficacy and a high degree of activity against the model Gram-positive bacterium, *S. aureus*, with an MIC close to that of temporin-1Ec (MIC 8µM) (23). However temporin-1Ee, which has an amino acid sequence quite similar to that of temporin-1Ec with just one chemically-conservative substitution (Val8 instead of Leu8 – Table 2.), is able to inhibit the growth of the model Gram-negative bacterium, *Escherichia coli*. Thus even conservative, single-site substitutions can dramatically change the spectrum of activity of a given temporin. The temporins and their analogues are thus well accepted candidates for the design of new antimicrobial peptides for reasons such as above, where small changes in amino acid sequence can result in dramatic improvements in antimicrobial activity but retaining a weak haemolytic activity (24-27). These factors bode well for translating temporins into clinically useful anti-infective agents to combat antibiotic-resistant pathogens.
Acknowledgments
This work was supported by National Natural Science Foundation, China (Grant No. 81402842), Natural Science Foundation of Fujian Province, China (Grant No. 2015J05162), Natural Science Foundation of Fujian Province, China (Grant No. 2016J01374), Fujian Provincial outstanding talents for Education and Scientific research (Grant No. JK2014016) and The Doctor Startup Foundation of Fujian Medical University (Grant No. 2012bs002).

Conflict of Interest statement
The authors declare that they have no conflict of interest.

Ethical statement
All applicable international, national, and/or institutional guidelines for the care and use of animals were followed.
Data deposition footnotes

The nucleotide sequence of the temporin-1Ee and temporine-1Re precursors from the skin secretion of the European edible frog, *Pelophylax kl.esculentus*, have been deposited in the EMBL Nucleotide Sequence Database under the accession code KT437658 and KT437659.
References


Legends to Figures

Figure 1. Nucleotide sequences of cloned cDNAs encoding the biosynthetic precursors of temporin-1Ee (A) and -1Re (B) and translated amino acid sequences of their open-reading frames. Putative signal peptides are double-underlined, mature peptides are single-underlined and the stop codons are indicated by asterisks. The G residue (italics) at the C-terminal of the mature peptide sequence serves as an amide donor after removal of the Lys residue (italics) with a carboxypeptidase. Residues G or GK (italics) at the end of the mature peptide sequence serve for C-terminal amidation.

Figure 2. (A) Alignment of full-length nucleotide sequences of clones encoding the precursors of temporin-1Ee and temporin-1Re. DNA sequences were copied in FASTA format, which were then subjected to multiple sequence alignment by the CLUSTALW program. (B) Alignment of open-reading frame amino acid sequences of preprotemporin-1Ee and preprotemporin-1Re. The conserved residues are shaded. Amino acid sequences were copied in FASTA format, which were then subjected to multiple sequence alignment by the CLUSTALW program.

Figure 3. Reverse phase HPLC chromatogram of the skin secretion from Rana esculenta with arrows indicating elution positions/retention times of the two novel antimicrobial peptides, temporin-1Re and temporin-1Ee. The Y-axis indicates absorbance units at λ=214nm.

Figure 4. (A) MALDI–TOF mass spectrum showing major peptide molecular ion in reverse phase HPLC fraction \( \sim 164 \) of Pelophylax kl. esculentus skin secretion corresponding in mass to temporin-1Ee (1402.08Da). (B) MALDI–TOF mass spectrum of peptide molecular ion in reverse phase HPLC fraction \( \sim 143 \) of skin extract corresponding to temporin-1Re (1012.32Da).
Figure 1

(A)

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1  ATGTTCCACA TGAAGAAAATC CCGTTTACCT CTTTTCTCTCC TTGGGACCAT
   TACAAGTGTT ACTTTCTTAG GGCAATGAG GAAAAAGAGG AACCTTGGTA
   NLSLCEERDAEEEEER
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   DPDPEERDVEMEKRFPLPV
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   TAGVLSRLFGR
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   TAAGCACTCT AAAACTCTTCT TGAACAGGCG TTTATTGGGT TTTTACACT
201  AACTTGGGAA ATAGAATGAG AATCTATCTG ATGTGGAATA TCAATTAGCT
   TTTGAACCTT TATCTTAAAC TTGTAGAGAC TACACCTTAT AGAATAATGCA
251  AAATGCAAGA CAGATGCTCT ATAAAAAAAT AAAGATATAA CATATAAAAAA
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(B)

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Figure 2

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Figure 3
Figure 4

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(B)