Analysis and Prediction of the Critical Regions of Antimicrobial Peptides Based on Conditional Random Fields

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Abstract

Antimicrobial peptides (AMPs) are potent drug candidates against microbes such as bacteria, fungi, parasites, and viruses. The size of AMPs ranges from less than ten to hundreds of amino acids. Often only a few amino acids or the critical regions of antimicrobial proteins matter the functionality. Accurately predicting the AMP critical regions could benefit the experimental designs. However, no extensive analyses have been done specifically on the AMP critical regions and computational modeling on them is either non-existent or settled to other problems. With a focus on the AMP critical regions, we thus develop a computational model AMPcore by introducing a state-of-the-art machine learning method, conditional random fields. We generate a comprehensive dataset of 798 AMP cores and a low similarity dataset of 510 representative AMP cores. AMPcore could reach a maximal accuracy of 90% and 0.79 Matthew's correlation coefficient on the comprehensive dataset and a maximal accuracy of 83% and 0.66 MCC on the low similarity dataset. Our analyses of AMP cores follow what we know about AMPs: High in glycine and lysine, but low in aspartic acid, glutamic acid, and methionine; the abundance of α-helical structures; the dominance of positive net charges; the peculiarity of amphipathicity. Two amphipathic sequence motifs within the AMP cores, an amphipathic α-helix and an amphipathic π-helix, are revealed. In addition, a short sequence motif at the N-terminal boundary of AMP cores is reported for the first time: arginine at the P(-1) coupling with glycine at the P1 of AMP cores occurs the most, which might link to microbial cell adhesion.

Introduction

Antimicrobial peptides (AMPs) are potent drug candidates against microbial organisms such as bacteria, fungi, parasites, and viruses. AMPs, which play an important role in innate immune responses to microbes, are derived from a broad spectrum of species such as archaea, bacteria, plants, and animals [1, 2]. Lysozyme is the first AMP discovered in human tissues and body fluids about 90 years ago [3]; α-purothionins found in wheat is the first AMP of plants [4]; gramicidin, a mixture of several AMPs, is one of the earliest AMPs derived from bacteria, Bacillus brevis [5]. However, the discovery of penicillin, the super antibiotic, might have hindered the
development of AMP researches in the mid-20th century. Not until 1980’s, the AMP researches revived. Then more AMPs were found including cecropins of silk moths [6], sarcotoxins of flies [7], defensins of rabbit macrophages [8], and magainins of frogs [9]. Up to date, thousands of AMPs are available and more than a dozen AMPs have entered clinical trials [10].

AMPs with various sizes have been documented in the literature. They range from ten to hundreds of amino acids. It is common to see that some AMPs would nest on other longer sequences. Such findings attribute antimicrobial effects to certain key active segments or critical regions of AMPs. Here we list three well-studied cases as examples: (I). Human LL-37 of human cathelicidin antimicrobial peptide (CAMP) expressed in different cells such as neutrophils, mast cells, epithelial cells, and macrophages [11–14]. Human LL-37 which consists of 37 amino acids beginning with double leucine is known to be antimicrobial [15]. In addition, KR-12 with only 12 amino acids is the shortest active segment of human LL-37 (18–29) [16]. (II). Human lactotransferrin, also known as human lactoferrin (hLF), found in various body fluids such as milk, tears, saliva, and nasal mucus as well as neutrophils [17–19]. Both hLF and its short helical segment with 11 amino acids hLF(21–31) demonstrated potent antimicrobial activities [20]. Similar cases were also seen in non-human LFs and their short segments [21, 22]. (III). Histidine-rich human histatin secreted from parotid and submandibular salivary glands [23]. Human histatin 8 with 12 amino acids is the shortest antimicrobial segment derived from histatin, which is common to most of the histatin family including histatin 3, 4, 5, 6, 7, 9, and 10 [24].

How AMPs defeat microbes have attracted researchers’ attention. It is believed that AMPs, abundant in cationic residues like lysine and arginine and scarce in anionic residues like aspartic acid and glutamic acid, interact with the anionic membranes of microbes to form transmembrane pores, thus causing abruption of microbes [10]. Besides, AMPs may enhance the progress of phagocytosis or the recruitment of leukocytes [25], and could even alter the gene expression of microbes [26].

Several computational methods have been applied to predict AMPs. They include quantitative matrix, discriminant analysis, artificial neural network, neuro-fuzzy interference, hidden Markov model, support vector machine, random forest, quantitative structure-activity relationship, and feature selection method [2, 27–32]. Generally speaking, these models utilized various features such as amino acid composition, protein secondary structure, net charge, and peptide aggregation to predict whether protein sequences are AMPs.

However, little is done in computationally identifying the critical regions of AMPs. To explain the differences between predicting AMPs and predicting AMP critical regions, the analogy of gene prediction is used. The current AMP prediction is like predicting whether genomic DNA sequences contain genes, but not where genes are located. Identifying the critical regions of AMPs is like finding which regions of DNA sequences encode genes. Modeling AMP critical regions requires understanding not only the differences between AMPs and non-AMPs but also how critical and non-critical AMPs regions are transited. Here we use the AMP critical regions defined as the shortest segments of the nested AMP families retaining antimicrobial effects deduced from current experimental evidence to model AMP cores.

Among these computational approaches, the AMPA server developed an unconventional method using high-throughput substitution data of 12-mer bactenecin against Pseudomonas aeruginosa to identify the active AMP stretches [33]. This method assigned each amino acid a bactericidal propensity value, which is the average of the bacterial half-maximal inhibitory concentration (IC50) of all the 12 variants of bactenecin [34]. According to the bactericidal propensity values, arginine and lysine, the basic residues, were the most lethal amino acids against microbes; aspartic acid and glutamic acid, the acid residues, were the least lethal.
To our knowledge, this is the first study to systematically extract the critical regions of AMPs or the cores of AMPs, examine their properties, and employ conditional random fields (CRFs) with multiple features to model them. Several important features of the AMP cores are investigated, including amino acid composition, protein secondary structures, net charges, amphipathicity, conserved protein domains, gapless alignments to highly similar protein sequences, AMPA bactericidal propensity, and peptide aggregation.

**Materials and Methods**

The framework of AMPcore is shown in **Fig. 1**. Nested AMP families were first generated. Next, the critical and non-critical regions of AMPs along with their features were determined. Our models based on these features were then trained using CRFs and were evaluated by 10-fold cross-validation.

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**Fig 1. Flowchart of AMPcore.** 2,497 non-redundant experimentally validated AMPs were obtained from CAMP release 2. 798 AMP nested families are recognized through self-comparison and their source proteins from NCBI nr database. The critical regions or cores of the nested families are then determined along with their features such as protein secondary structures and conserved domains. AMPcore are built using CRFs based on the nested AMPs and the performances are examined using 10-fold cross-validation.

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Data collection of AMP critical regions

AMP critical regions are defined here as the shortest experimentally-validated segments of AMP proteins retaining antimicrobial function. In computational terms, AMP critical regions are the minimal substrings of AMP protein sequences supported by experimental evidence. It should be noted that shorter AMP cores which have not yet been found by experiments might exist.

The AMP critical regions in this study were determined by either nested AMP families or source proteins. 2,497 experimentally validated non-redundant AMPs were obtained from CAMP release 2 [2]. 158 nested AMP families were directly extracted from the experimentally validated dataset. In addition, each AMP sequence in the dataset was run against the NCBI non-redundant (nr) protein database using BLAST to search for its full-length source protein. The longer source proteins were also included in the nested families. Each nested AMP family thus contains at least one short AMP sequence and at least one AMP source protein. Those families with single members without valid source proteins, were not considered. 798 nested AMP families were collected in our final collection. Based on this collection, 510 representative nested AMP families were also generated by filtering out highly similar families with over 70% sequence identity using CD-HIT [35]. Such cutoff threshold was chosen by following Wang et al. [27].

The AMP critical regions of a nested AMP family can be described as follows: Let $P_m = P_{m_1} P_{m_2} \ldots P_{m_n}$ be a protein sequence with $n$ residues. Given a nested AMP family $P = \{P_1, \ldots, P_k\}$ with $k \geq 2$ and $|P_i| = n_i$, the AMP critical region is determined by the shortest AMP $P_s$.

$$P_s := P_s \in P \land P_s \subseteq \bigcap_{i=1}^{k} P_i \land |P_s| = \min_{1 \leq i \leq k} (|P_i|)$$

where $P_s$ is a member of the family and also a substring of all the other AMP sequences in the family. In our study, $40 \geq |P_s| \geq 7$. Outside $P_s$ is considered to be non-critical for antimicrobial function.

The non-critical regions of a nested AMP family are determined once $P_s$ is found. The longest sequence in the family marked with the critical and non-critical regions is utilized to train our model. Fig. 2 illustrates the concept of a nested AMP family.

**Conditional random fields**

CRFs are a discriminative undirected probabilistic graphical model proposed by Lafferty et al. [36]. Both generative and discriminative models are probabilistic models. However, generative models such as hidden Markov models (HMMs) are based on the joint probability distribution $P(Y, X)$ and discriminative models such as maximum entropy Markov models (MEMMs) and CRFs are based on the conditional probability distribution $P(Y | X)$, where variables $X$ and $Y$

\[ P_A: \quad \begin{array}{cccccccccccc}
\bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet \\
\end{array} \\
\text{P}_B: \quad \begin{array}{cccccccccccc}
\bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet \\
\end{array} \\
\text{P}_C: \quad \begin{array}{cccccccccccc}
\bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet & \bullet \\
\end{array} \\
\]

**Fig 2. A simple diagram of a nested AMP family.** This nested family has three AMPs: $P_A$, $P_B$, and $P_C$. Each circle represents an amino acid. The circles in gray are the critical regions of the family, which are the overlapping residues shared by all the family members; those in white are the non-critical regions. Note: here a family has at least two members and one member with at least seven amino acids long would map to the critical region entirely.

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represent observations and labels respectively. The fundamental differences require generative models to model \(P(X)\), which is difficult when \(P(X)\) involves multiple attributes. Generally speaking, CRFs outperform traditional generative graphical models like HMMs because they can relax the strong independence assumptions made in generative models [36].

In addition, CRFs overcome the label bias problem which other discriminative models based on next-state classifiers like MEMMs suffer. Given a current state, these models determine which state to go next by comparing all outgoing transitions of the current state, not all transitions of the models. The per-state computation is biased in favor of the states with fewer outgoing transitions. Instead CRFs are designed to consider the entire label sequence given the observation sequence in terms of conditional probability.

We would like to address our problem as a sequence labeling problem. A typical sequence labeling problem is to predict a sequence of categorical labels \(Y = (y_1, y_2, \ldots, y_n)\) given a sequence of observations \(X = (x_1, x_2, \ldots, x_i)\). CRFs are known to excel in sequence labeling tasks such as part-of-speech tagging and text segmentation in natural language sequences [36, 37], image segmentation in video sequences [38], and gene finding [39], RNA secondary structural alignment [40], protein structural instability [41] and protein domain boundary in biological sequences [42].

To determine the most probable sequence of labels \(\hat{Y}\), that is,

\[
\hat{Y} = \arg \max_{Y} P(Y|X)
\]

CRFs use an exponential function to compute the conditional probability of labels with respect to observations as follows:

\[
P(Y|X) = \frac{\sum_{i=1}^{n} \sum_{j} \lambda_j f_j(y_{i-1}, y_i, x, i)}{Z(X)}
\]

where index \(i\) represents position \(i\) in the sequence, \(j\) represents the \(j\)th observed feature, \(\lambda_j\) is a \(j\)th weight vector, and \(f_j\) is a \(j\)th feature vector which considers both a transition feature \(t_j(y_{i-1}, y_i, x, i)\) between position \(i-1\) and position \(i\) and a state feature \(s_j(y_i, x, i)\) at position \(i\) in the label sequence given the observation \(x\). \(Z(X)\) is an observation-dependent global normalization function which enumerates all possible labels given the observation \(x\) as follows:

\[
Z(X) = \sum_j e^{\sum_{i=1}^{n} \sum_{j} \lambda_j f_j(y_{i-1}, y_i, x, i)}
\]

To maximize the likelihood of \(\hat{Y}\), the optimal \(\lambda\) is found using L-BFGS algorithm [43], a limited-memory quasi-Newton method, in the iterative training stage. CRFs guarantee to reach a global maximum likelihood, for they are globally conditioned on the observation \(x\). More detail about the parameters in CRFs can be found in Lafferty et al. [36]. In our case, the conditional probability distribution treats protein sequences as the primary observations and critical/non-critical regions as the labels. Each residue of the protein sequence would be mapped to an associated label.

In this study, CRF++ version 0.58 was utilized [44]. The rules for training our CRF models were obtained from the template files.
Observed Features

We examined eight different aspects of the critical regions of AMPs:

1. Protein primary structures
2. Protein secondary structures
3. Conserved protein domains
4. Short pairwise alignments
5. AMPA
6. Aggregation
7. Net charges
8. Amphipathicity

A sliding window was applied to study the features/observations of the critical and non-critical regions of AMPs. The size of the sliding window would affect the testing results. If the window size gets too large, excessive parameters need to be trained, which may add much extraneous information; if the window size is too small, it may not include enough essential information. The window size was selected to be five in this study. Empirically we found that such size was an appropriate choice for the model to have a solid performance.

Protein secondary structures. Protein secondary structure prediction (PSSpred version 2.0), a neural network classifier taken from the famous I-TASSER server, was utilized to predict the secondary structure of a peptide [45]. Each amino acid in an AMP was classified into α-helix, β-sheet, or random coil.

Conserved protein domains. Pfam domains based on multiple sequence alignments [46] were selected to represent known conserved protein domains. Each sequence was checked against Pfam. The regions mapped to the protein domains were labeled as conserved. Otherwise, not conserved.

Short Pairwise alignments. We designed an alignment procedure to search against the AMP core database. Neither global sequence alignment nor local sequence alignment was utilized, for this type of alignment focuses on long similar hits. Instead a gapless alignment using BLOSUM62 matrix was performed to search shorter similar hits, where only positive and neutral amino acid substitution were allowed. In addition, a heuristic approach by standardizing the alignment score by sequence length was used to rank the hits.

AMPA. AMPA was utilized to locate AMP stretches [33]. The default parameters were used to determine whether each amino acid resided in antimicrobial domains or not.

Aggregation. AGGRESCAN was utilized to estimate the aggregation tendencies of a peptide [47]. AGGRESCAN, which applied aggregation propensities of amino acids derived from the experimental data of β-amyloid peptides, was a good indicator of in vivo aggregation.

Net charges. A N-mer sliding window was utilized to go through the sequence. There were three categories of residues: leading, positive, and non-positive. All of the beginning N-1 residues were leading. The net charge of the N-mer is the difference between the count of the positive charged residues and that of the negative charged residues. Here N was set to 7.

Amphipathicity. Amphipathic moment, also known as the mean hydrophobic moment, was utilized to measure the amphipathicity of a peptide (S1 Equation) [48]. The amphipathicity of a peptide is that along the internal axis of the peptide backbone, one side contains non-polar residues and the other side contains polar or charged residues. Amphipathic moment quantifies such character by calculating the average hydrophobic difference of the two sides along the
axis. HMOMENT with the default parameters from the EMBOSS version 6.5.7 was used to determine whether the peptide segment is amphipathic [49].

**Evaluation**

The following statistical measures were utilized to evaluate model performance. They are sensitivity, specificity, accuracy, and Matthew’s coefficient of correlation (MCC) defined as follows:

\[
\text{Sensitivity} = \frac{TP}{TP + FN}
\]

\[
\text{Specificity} = \frac{TN}{FP + TN}
\]

\[
\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN}
\]

\[
\text{MCC} = \frac{TP \times TN - FP \times FN}{\sqrt{(TP + FP)(TP + FN)(TN + FP)(TN + FN)}}
\]

TP, TN, FP, and FN are true positives, true negatives, false positives, and false negatives, respectively. Sensitivity is called the true positive rate; specificity is also known as the true negative rate. Accuracy is the percentage of the correct predictions among all the positive and negative data. MCC, which could range from −1 to 1, is a good performance indicator for a binary classifier. The larger the MCC value, the better the classifier.

**Results**

The comparisons of the AMP cores and source proteins, which consist of the AMP critical and non-critical regions, were performed. The performance of our CRF models were then evaluated.

**Amino acid composition of the critical regions of AMPs**

The comparison of amino acid composition of the critical regions of AMPs was shown in Fig. 3. Compared to the source proteins, the AMP cores showed higher percentages of glycine and lysine, but lower percentages of aspartic acid, glutamic acid, and methionine using the means and medians. The results by the means and medians were consistent except for arginine. For arginine, the AMP cores had a higher mean but much lower median than the source proteins. Interestingly, glycine, the smallest but flexible residue, was the most abundant residue in the critical regions of AMPs. In addition, the AMP cores and the source proteins shared similar hydrophobic contents. The difference of the total percentage of the hydrophobic residues between the AMP cores and the source proteins was negligible although about two-fifths of the overall residues were hydrophobic.

**Secondary structures of the critical regions of AMPs**

The statistical analysis of protein secondary structures of the AMP cores was performed as shown in Fig. 4. The protein secondary structures were classified into α-helix, β-strand, or coil. More than half of AMP cores were α-helix, which was higher than the source proteins. In addition, the AMP cores had lower tendency to be coil than the source proteins. β-strand structures occurred infrequently in both the AMP cores and source proteins.
Net charges of the critical regions of AMPs

The net-charge distributions of the AMP cores and the source proteins were compared in Fig. 5. The net-charge plot of AMP cores is similar to that of AMPs albeit with some slight differences [50]. The AMP cores predominantly had a positive net charge. About half of the AMP cores had a net charge between +2 and +4 and less than 5% of the AMP cores had a negative net charge. Compared to the AMP cores, the source proteins did not display a strong net-charge preference. Contrarily over one third of the source proteins had a negative net charge.

Amphipathicity of the critical regions of AMPs

Fig. 6 compared the amphipathicity distributions of the AMP cores and the source proteins in terms of α helixes. The result showed that the AMP cores tended to have higher amphipathic values than the source proteins, suggesting that the AMP cores fit amphipathic α-helical structures better. However, the two amphipathicity distributions were less sufficiently separated than the two net-charge distributions in Fig. 5.
Sequence motifs of the critical regions of AMPs

The sequence motifs of the AMP cores were examined using MEME version 4.9.1 with the default parameters [51]. Three motifs were obtained as shown in Fig. 7. Two of the three motifs showed periodic occurrences of positively charged residues and the other was a cysteine-based motif. Both of the periodically charged motifs were further examined on a helical wheel plot. Interestingly they demonstrated a clear amphipathic property while one suited better into an amphipathic π-helical structure (S1 Fig.) and the other was a fine amphipathic α helix with one hydrophobic side and one hydrophilic side dominated with the positive charges (S2 Fig.). The cysteine-based motif acted like a hinge. The conserved cysteines at the edges of the cysteine-based motif were found to involve in the formation of disulfide bridges, not to each other, but to other distant cysteines; the glycine at the P9 could make the hinge motif more flexible.

Boundaries of the critical regions of AMPs

The N- and C-terminal boundaries of the AMP cores were examined as shown in Fig. 8. Just outside the N-terminal ends of the AMP cores (P-1), a weak preference for charged residues such as arginine was observed. Our analysis further showed the boundary arginine frequently paired with glycine, alanine, serine, and phenylalanine at the N-terminal end of the AMP cores. Particularly, arginine at the P-1 with glycine at the P1 occurred the most (S3 Fig.). In addition, adjacent to the C-terminal end of the AMP cores, glycine and positively charged residues were preferred. However, we found that the boundary bias at the C-terminal AMP cores was less apparent. More details could be found at S4 Fig.
Performance comparison of CRF models

To evaluate the importance of each feature, the feature ablation studies were performed, which removes one of the testing features at a time. All the CRF models in the ablation studies were examined using 10-fold cross-validation on the testing datasets. Table 1 and Table 2 showed the performance comparison of the CRF models on the datasets of 798 AMP cores and 510 representative AMP cores respectively.

Not all of our testing features contribute to the model performance. Table 1 and Table 2 indicate that among these testing features, amino acid sequence, protein secondary structure, net charge, and amphipathicity were vital. However, aggregation, gapless alignment, and AMPA seemed less important, for they improved the CRF models little or none. Our CRF models could reach a maximal 90% accuracy and 0.79 MCC on the dataset of 798 AMP cores, but
dropped to a maximal 83% accuracy and 0.66 MCC on the dataset of 510 representative AMP cores.

Performance comparison of the AMP prediction models in predicting AMP cores

To verify if predicting AMP cores could be simply substituted by predicting AMPs, several AMP prediction models from CAMP webserver were investigated, including SVM, random forest, artificial neural network, and discriminant analysis [29]. Unlike our AMPcore models using these eight features built under 10-fold cross-validation, all the CAMP models were built on the full set of CAMP data with all of the AMP cores. A 10-mer sliding window, the minimum window size allowed in CAMP, was then utilized for these AMP models to go through the test cases of the AMP cores. As long as the predictors assigned the peptide segments to be AMPs, they would be marked as the critical regions. On the comprehensive dataset, the best CAMP models could achieve a maximal accuracy of 70% and 0.40 MCC; on the low similarity dataset, they could reach a maximal accuracy of 68% and 0.37 MCC (Table 3 and Table 4).

Table 1. Feature ablation study of AMPcore on the comprehensive dataset using 10-fold cross-validation.

| Features            | Sensitivity | Specificity | Accuracy | MCC  |
|---------------------|-------------|-------------|----------|------|
| All                 | 89.1        | 90.1        | 89.6     | 0.79 |
| Primary structures  | 73.5        | 88.0        | 80.8     | 0.62 |
| Secondary structures| 88.4        | 89.1        | 88.8     | 0.78 |
| Net charges         | 88.9        | 88.9        | 88.9     | 0.78 |
| Amphipathicity      | 88.3        | 90.0        | 89.1     | 0.78 |
| Conserved domains   | 88.4        | 89.5        | 88.9     | 0.78 |
| Aggregation         | 88.9        | 89.3        | 89.1     | 0.78 |
| Gapless alignments  | 89.3        | 89.6        | 89.5     | 0.79 |
| AMPA                | 89.0        | 90.1        | 89.6     | 0.79 |

The symbol ‘−’ stands for “subtracting”.

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Discussion

This was the first study to systematically extract, examine, and model the critical regions of AMPs. Several properties of the AMP cores were found to be common with those of the AMPs, but the boundary bias of the AMP cores were newly discovered in this study. We demonstrated that the general-purposed AMP prediction tools were not suited for the prediction of AMP cores and our ad hoc CRF model with multiple features would work better.

Several key features are common to both AMPs and AMP cores. The AMP cores are abundant in glycine and lysine, but deficient in methionine and acidic residues, the same as the AMPs [1]. The N-terminal ends of the AMP cores are relatively rich in glycine, similar to the findings in antibacterial peptides [28]. Besides, as cationic AMPs prevail, the AMP cores are predominantly having a positive net charge. Further, the periodic occurrences of positively charged residues lead to the amphipathicity of the AMP cores, also known to the AMPs [10]. Like what have been suggested in AMPs, these properties might help the AMP cores to interact with anionic microbial cell membranes.

There are also novel findings reporting for the first time. Plenty of charged residues were found right outside the N-terminal and C-terminal ends of the AMP cores. These charged residues might help stabilize the AMPs attached to microbes. In addition, we discovered that R/G, R/F, R/S, and R/A at the boundary of the N-terminal AMP cores occurred more frequently than would be expected by chance. Particularly, R/G was the most apparent.

Why arginine at the P-1 of the source proteins coupling with glycine at the P1 of the N-terminal AMP cores is not clear. While AMP cores remain antimicrobial activities, the boundaries of AMP cores might provide accessory functions. Two possible explanations are proposed: (1) The R/G site might be a cell adhesion signal to microbes, similar to the RGD peptides of fibronectin, an adhesion protein, to cell hosts [52]. Fibronectin-binding protein might be one of the

**Table 2. Features ablation study of AMPcore on the low similarity dataset using 10-fold cross-validation.**

| Features          | Sensitivity | Specificity | Accuracy | MCC  |
|-------------------|-------------|-------------|----------|------|
| All               | 82.3        | 84.0        | 83.2     | 0.66 |
| - Primary structures | 65.3        | 82.0        | 73.7     | 0.48 |
| - Secondary structures | 81.2        | 82.0        | 81.6     | 0.63 |
| - Net charges     | 81.7        | 82.5        | 81.1     | 0.64 |
| - Amphipathicity  | 81.9        | 82.2        | 82.0     | 0.64 |
| - Conserved domains | 81.9        | 82.5        | 82.2     | 0.65 |
| - Aggregation     | 82.2        | 83.6        | 82.9     | 0.66 |
| - Gapless alignments | 82.4        | 84.0        | 83.2     | 0.66 |
| - AMPA            | 81.3        | 84.3        | 82.8     | 0.66 |

The symbol ‘-’ stands for "subtracting".

**Table 3. Performance comparison with AMP prediction models on the comprehensive dataset of 798 AMP cores.**

| Model    | Sensitivity | Specificity | Accuracy | MCC  |
|----------|-------------|-------------|----------|------|
| CAMP.SVM | 55.0        | 59.8        | 56.7     | 0.14 |
| CAMP.RF  | 47.0        | 47.9        | 47.5     | -0.05|
| CAMP.ANN | 62.7        | 82.2        | 68.4     | 0.41 |
| CAMP.DA  | 71.6        | 68.2        | 69.7     | 0.40 |
| AMPcore  | 89.1        | 90.1        | 89.6     | 0.79 |

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receptors for the naturally occurring AMPs. (2) The R/G site might involve with the interaction of RNA, for RG-rich protein domains are known to affect RNA binding [53]. Additional studies, which are beyond the scope of this study, are still required to determine the role of R/G at the boundary of the N-terminal AMP cores.

Other than the boundary preferences, the secondary structures of the AMP cores were quantified. Most of the AMP cores were α-helix, a common structure for protein-protein interactions, which was similar to our previous finding in antiviral peptides [54]. Besides, the AMP cores possessed a stronger amphipathic helical character than the background. Two amphipathic sequence motifs within the AMP cores were found: One α-helix and one π-helix. A previous study even suggested that relatively infrequent π-helix was directly linked to the active sites of proteins [55]. Interestingly, our findings of the AMP cores supported this assertion.

Predicting AMP cores could not be replaced simply by predicting AMPs. Table 3 and Table 4 demonstrate that by far our CRF models of AMP cores outperformed these current AMP prediction models. CRFs were chosen to build AMPcore, for they are generally superior in modeling the sequential data [36]. In fact, in order to reach optimal performances in predicting AMP cores, our CRF model required not only amino acid sequences but also additional features such as protein secondary structures, net charges, and amphipathicity. Accurately predicting AMP cores in protein sequences would facilitate experimental designs.

Supporting Information

S1 Fig. Amphipathic π-helical wheel plot of an AMP core containing motif I. (TIF)

S2 Fig. Amphipathic α-helical wheel plot of an AMP core containing motif III. (TIF)

S3 Fig. Relative distribution of amino acid coupling at the N-terminal boundary of AMP cores. Each row represents the residue at the P(-1) of the boundary of the source protein; Each column represents the residue at the P1 of the N-terminal AMP cores. Each cell represents the log-odd value for such pattern against the background. A heat map is used to scale the log-odd value: low (black) to high (white). (TIF)

S4 Fig. Relative distribution of amino acid coupling at the C-terminal boundary of AMP cores. Each row represents the residue at the P(-1) of the C-terminal AMP cores; Each column represents the residue at the P1 of the boundary of the source proteins. Each cell represents the log-odd value for such pattern against the background. A heat map is used to scale the log-odd value: low (black) to high (white). (TIF)

Table 4. Performance comparison with AMP prediction models on the low similarity dataset of 510 AMP cores.

| Model   | Sensitivity | Specificity | Accuracy | MCC  |
|---------|-------------|-------------|----------|------|
| CAMP.SVM | 54.9        | 60.1        | 56.8     | 0.15 |
| CAMP.RF  | 47.4        | 49.0        | 48.2     | -0.04|
| CAMP.ANN | 60.5        | 78.9        | 65.7     | 0.35 |
| CAMP.DA  | 70.1        | 67.0        | 68.4     | 0.37 |
| AMPcore  | 82.3        | 84.0        | 83.2     | 0.66 |

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Author Contributions
Conceived and designed the experiments: KYC TPL CKW. Performed the experiments: KYC TPL LYS CKW. Analyzed the data: KYC TPL. Contributed reagents/materials/analysis tools: KYC. Wrote the paper: KYC.

References
1. Wang G, Li X, Wang Z. APD2: the updated antimicrobial peptide database and its application in peptide design. Nucleic acids research. 2009; 37(Database issue):D933–7. doi:10.1093/nar/gkn823 PMID: 18957441
2. Waghu FH, Gopi L, Barai RS, Ramteke P, Nizami B, Idicula-Thomas S. CAMP: Collection of sequences and structures of antimicrobial peptides. Nucleic acids research. 2014; 42(Database issue):D1154–8. doi: 10.1093/nar/gkt1157 PMID: 24265220
3. Fleming A. On a remarkable bacteriolytic element found in tissues and secretions. Proc R Soc London B. 1922; 93:306–17.
4. Stuart LS, Harris TH. Bactericidal and fungicidal properties of a crystalline protein isolated from un-bleached wheat flour. Cereal Chem. 1942; 19:288–300.
5. Dubos RJ. Studies on a bactericidal agent extracted from a soil bacillus: I. Preparation of the agent. Its activity in vitro. J Exp Med. 1939; 70:1–10. PMID: 19870884
6. Steiner H, Hultmark D, Engstrom A, Bennich H, Boman HG. Sequence and specificity of two antibacterial proteins involved in insect immunity. Nature. 1981; 292(5820):246–8. PMID: 7019715
7. Okada M, Natori S. Purification and characterization of an antibacterial protein from haemolymph of Sarcophaga peregrina (flesh-fly) larvae. The Biochemical journal. 1983; 211(3):727–34. PMID: 6892366
8. Patterson-Delafield J, Szklarek D, Martinez RJ, Lehrer RI. Microbicidal cationic proteins of rabbit alveolar macrophages: amino acid composition and functional attributes. Infection and immunity. 1981; 31(2):723–31. PMID: 7216471
9. Zasloff M. Magainins, a class of antimicrobial peptides from Xenopus skin: isolation, characterization of two active forms, and partial cDNA sequence of a precursor. Proceedings of the National Academy of Sciences of the United States of America. 1987; 84(15):5449–53. PMID: 3299384
10. Fjell CD, Hiss JA, Hancock RE, Schneider G. Designing antimicrobial peptides: form follows function. Nature reviews Drug discovery. 2012; 11(1):37–51. doi: 10.1038/nrd3591 PMID: 22173434
11. Cowland JB, Johnsen AH, Borregaard N. hCAP-18, a cathelin/pro-bactenecin-like protein of human neutrophil specific granules. FEBS letters. 1995; 368(1):173–6. PMID: 7615076
12. Di Nardo A, Vitiello A, Gallo RL. Cutting edge: mast cell antimicrobial activity is mediated by expression of cathelicidin antimicrobial peptide. J Immunol. 2003; 170(5):2274–8. PMID: 12594247
13. Murakami M, Lopez-Garcia B, Braff M, Dorschner RA, Gallo RL. Postsecretory processing generates multiple cathelicidins for enhanced topical antimicrobial defense. J Immunol. 2004; 172(5):3070–7. PMID: 14978112
14. Yuk JM, Shin DM, Lee HM, Yang CS, Jin HS, Kim KK, et al. Vitamin D3 induces autophagy in human monocytes/macrophages via cathelicidin. Cell host & microbe. 2009; 6(3):231–43.
15. Sorensen OE, Follin P, Johnsen AH, Calafat J, Tjabringa GS, Hiemstra PS, et al. Human cathelicidin, hCAP-18, is processed to the antimicrobial peptide LL-37 by extracellular cleavage with proteinase 3. Blood. 2001; 97(12):3951–9. PMID: 11390939
16. Wang G. Structures of human host defense cathelicidin LL-37 and its smallest antimicrobial peptide KR-12 in lipid micelles. The Journal of biological chemistry. 2008; 283(47):32637–43. doi: 10.1074/jbc.M805533200 PMID: 18816205
17. Masson PL, Heremans JF. Lactoferrin in milk from different species. Comparative biochemistry and physiology B, Comparative biochemistry. 1971; 39(1):119–29. PMID: 4998849

18. Rado TA, Wei XP, Benz EJ Jr. Isolation of lactoferrin cDNA from a human myeloid library and expression of mRNA during normal and leukemic myelopoiesis. Blood. 1987; 70(4):969–93. PMID: 3477300

19. Orsi N. The antimicrobial activity of lactoferrin: current status and perspectives. Biometals: an international journal on the role of metal ions in biology, biochemistry, and medicine. 2004; 17(3):189–96.

20. Chapple DS, Hussain R, Joannou CL, Hancock RE, Odell E, Evans RW, et al. Structure and association of human lactoferrin peptides with Escherichia coli lipopolysaccharide. Antimicrobial agents and chemotherapy. 2004; 48(6):2190–8. PMID: 15155221

21. Bellamy W, Takase M, Yamauchi K, Wakabayashi H, Kawase K, Tomita M. Identification of the bacterial domain of lactoferrin. Biochimica et biophysica acta. 1992; 1121(1–2):130–6.

22. Adao R, Nazmi K, Bolscher JG, Bastos M. C- and N-truncated antimicrobial peptides from LFampin 265–284: Biophysical versus microbiology results. Journal of pharmacy & bioallied sciences. 2011; 3(1):60–9.

23. Oppenheim FG, Xu T, McMillian FM, Levitz SM, Offner GD, et al. Histatins, a novel family of histidine-rich proteins in human parotid secretion. Isolation, characterization, primary structure, and fungistatic effects on Candida albicans. The Journal of biological chemistry. 1988; 263(16):7472–7. PMID: 3286634

24. Yin A, Margolis HC, Grogan J, Yao Y, Troxler RF, Oppenheim FG. Physical parameters of hydroxyapatite adsorption and effect on candidacidal activity of histatins. Archives of oral biology. 2003; 48(5):361–8. PMID: 12711380

25. Hancock RE, Diamond G. The role of cationic antimicrobial peptides in innate host defences. Trends in microbiology. 2000; 8(9):402–10. PMID: 10989307

26. Hong RW, Shchepevtov M, Weiser JN, Axelsen PH. Transcriptional profile of the Escherichia coli response to the antimicrobial insect peptide cecropin A. Antimicrobial agents and chemotherapy. 2003; 47(1):1–6. PMID: 12499161

27. Wang P, Hu L, Liu G, Jiang N, Chen X, Xu J, et al. Prediction of antimicrobial peptides based on sequence alignment and feature selection methods. PloS one. 2011; 6(4):e18476. doi: 10.1371/journal.pone.0018476 PMID: 21533231

28. Lata S, Mishra NK, Raghava GP. AntiBP2: improved version of antibacterial peptide prediction. BMC bioinformatics. 2010; 11 Suppl 1:S19. doi: 10.1186/1471-2105-11-S1-S19 PMID: 20122190

29. Thomas S, Karnik S, Barai RS, Jayaraman VK, Idicula-Thomas S. CAMP: a useful resource for research on antimicrobial peptides. Nucleic acids research. 2010; 38(Database issue):D774–80. doi: 10.1093/nar/gkp1021 PMID: 19923233

30. Fernandes FC, Rigden DJ, Franco OL. Prediction of antimicrobial peptides based on the adaptive neuro-fuzzy inference system application. Biopolymers. 2012; 98(4):280–7. PMID: 23193592

31. Torrent M, Andreu D, Nogues VM, Boix E. Connecting peptide physicochemical and antimicrobial properties by a rational prediction model. PloS one. 2011; 6(2):e16968. doi: 10.1371/journal.pone.0016968 PMID: 21347392

32. Jenssen H, Fjell CD, Cherkasov A, Hancock RE. QSAR modeling and computer-aided design of antimicrobial peptides. Journal of peptide science: an official publication of the European Peptide Society. 2008; 14(1):110–4.

33. Torrem D, Di Tommaso P, Pulido D, Nogues MV, Notredame C, Boix E, et al. AMPA: an automated web server for prediction of protein antimicrobial regions. Bioinformatics. 2012; 28(1):130–1. doi: 10.1093/bioinformatics/btr604 PMID: 22053077

34. Hilpert K, Volkmer-Engert R, Walter T, Hancock RE. High-throughput generation of small antibacterial peptides with improved activity. Nature biotechnology. 2005; 23(8):1008–12. PMID: 16041386

35. Huang Y, Niu B, Gao Y, Fu L, Li W. CD-HIT Suite: a web server for clustering and comparing biological sequences. Bioinformatics. 2010; 26(5):680–2. doi: 10.1093/bioinformatics/btp003 PMID: 20053844

36. Sha F, Pereira F, editors. Shallow parsing with conditional random fields. Proceedings of the 2003 Conference of the North American Chapter of the Association for Computational Linguistics on Human Language Technology–Volume 1; 2003: Association for Computational Linguistics.

37. He X, Zemel RS, Carreira-Perpinnd M, editors. Multiscale conditional random fields for image labeling. Computer vision and pattern recognition, 2004 CVPR 2004 Proceedings of the 2004 IEEE computer society conference on; 2004: IEEE.
39. DeCaprio D, Vinson JP, Pearson MD, Montgomery P, Doherty M, Galagan JE. Conrad: gene prediction using conditional random fields. Genome research. 2007; 17(9):1389–98. PMID: 17690204
40. Sato K, Sakakibara Y. RNA secondary structural alignment with conditional random fields. Bioinformatics. 2005; 21 Suppl 2:i237–42. PMID: 16204111
41. Wang L, Sauer UH. OnD-CRF: predicting order and disorder in proteins using [corrected] conditional random fields. Bioinformatics. 2008; 24(11):1401–2. doi: 10.1093/bioinformatics/btn132 PMID: 18430742
42. Zhang XY, Lu LJ, Song Q, Yang QQ, Li DP, Sun JM, et al. DomHR: accurately identifying domain boundaries in proteins using a hinge region strategy. PloS one. 2013; 8(4):e60559. doi: 10.1371/journal.pone.0060559 PMID: 23593247
43. Liu DC, Nocedal J. On the limited memory BFGS method for large scale optimization. Mathematical programming. 1989; 45(1–3):503–28.
44. Kudo T. CRF++: Yet Another CRF toolkit (website). 2013 Available: http://crfpp.googlecode.com/svn/trunk/doc/index.html. Accessed 2014 April 7.
45. Zhang Y. PSSpred: A multiple neural network training program for secondary protein structure prediction (website). 2012 Available: http://zhanglab.ccmb.med.umich.edu/PSSpred. Accessed 2014 April 7.
46. Punta M, Coggill PC, Eberhardt RY, Mistry J, Tate J, Boursnell C, et al. The Pfam protein families database. Nucleic acids research. 2012; 40(Database issue):D290–301. doi: 10.1093/nar/gkr1065 PMID: 22127870
47. Conchillo-Sole O, de Groot NS, Aviles FX, Vendrell J, Daura X, Ventura S. AGGRESCAN: a server for the prediction and evaluation of "hot spots" of aggregation in polypeptides. BMC bioinformatics. 2007; 8:65. PMID: 17324296
48. Eisenberg D, Weiss RM, Terwilliger TC. The helical hydrophobic moment: a measure of the amphiphlicity of a helix. Nature. 1982; 299(5881):371–4. PMID: 7110359
49. Rice P, Longden I, Bleasby A. EMBOSS: the European Molecular Biology Open Software Suite. Trends in genetics: TIG. 2000; 16(6):276–7. PMID: 10827456
50. Wang G. A database view of naturally occurring antimicrobial peptides: nomenclature, classification and amino acid sequence analysis. Antimicrobial Peptides: discovery, design and novel therapeutic strategies. Advances in molecular and cellular microbiology. Cambridge, Massachusetts, USA: CABI; 2010.
51. Bailey TL, Boden M, Buske FA, Frith M, Grant CE, Clementi L, et al. MEME SUITE: tools for motif discovery and searching. Nucleic acids research. 2009; 37(Web Server issue):W202–8. doi: 10.1093/nar/gkp335 PMID: 19458158
52. Ruoslathi E, Pierschbacher MD. Arg-Gly-Asp: a versatile cell recognition signal. Cell. 1986; 44(4):517–8. PMID: 2418990
53. McBride AE, Conboy AK, Brown SP, Ariyachet C, Rutledge KL. Specific sequences within arginine-glycine-rich domains affect mRNA-binding protein function. Nucleic acids research. 2009; 37(13):4322–30. doi: 10.1093/nar/gkp349 PMID: 19454603
54. Chang KY, Yang JR. Analysis and prediction of highly effective antiviral peptides based on random forests. PloS one. 2013; 8(8):e70166. doi: 10.1371/journal.pone.0070166 PMID: 23940542
55. Weaver TM. The pi-helix translates structure into function. Protein science: a publication of the Protein Society. 2000; 9(1):201–6. PMID: 10739264