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StaRProtein, A Web Server for Prediction of the Stability of Repeat Proteins

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Abstract

Repeat proteins have become increasingly important due to their capability to bind to almost any proteins and the potential as alternative therapy to monoclonal antibodies. In the past decade repeat proteins have been designed to mediate specific protein-protein interactions. The tetratricopeptide and ankyrin repeat proteins are two classes of helical repeat proteins that form different binding pockets to accommodate various partners. It is important to understand the factors that define folding and stability of repeat proteins in order to prioritize the most stable designed repeat proteins to further explore their potential binding affinities. Here we developed distance-dependant statistical potentials using two classes of alpha-helical repeat proteins, tetratricopeptide and ankyrin repeat proteins respectively, and evaluated their efficiency in predicting the stability of repeat proteins. We demonstrated that the repeat-specific statistical potentials based on the two classes of repeat proteins showed paramount accuracy compared with non-specific statistical potentials in: 1) discriminate correct vs. incorrect models 2) rank the stability of designed repeat proteins. In particular, the statistical scores correlate closely with the equilibrium unfolding free energies of repeat proteins and therefore would serve as a novel tool in quickly prioritizing the designed repeat proteins with high stability. StaRProtein web server was developed for predicting the stability of repeat proteins.

Introduction

Repeat protein scaffolds are commonly found in all kingdoms of life. They typically function in mediating specific protein-protein interactions which are essential for various biological functions [1]. Repeat proteins are comprised of tandem arrays of short repeat motifs that stack together to form extended super-helical structure. So far more than twenty classes of repeat proteins have been identified, among which the most abundant are ankyrin repeat (AR), leucine-rich repeat (LRR), armadillo repeat (ARM), helical-repeat (HEAT) and tetrotricopeptide repeat (TPR) proteins.
Repeat proteins are attractive alternatives to antibodies due to their stability and ease of production as well as high binding affinities and specificity [2],[3]. In contrast to some repeat-containing proteins such as LRR and HEAT that bind a specific ligand with preferred secondary structure, TPR and AR proteins can bind with diverse proteins [4]. E.g. two discrete TPR domains in Hsp organizing protein (HOP) associate with molecular chaperone proteins Hsp70 and Hsp90, both being emerging cancer targets [5],[6],[7]. Envelope glyproteins, gp120 and gp41 mediate the entry of HIV-1 virus, and thus both are attractive anti-HIV targets [8]. Due to versatile binding profile of TPR and AR proteins, they can serve as useful scaffolds to mediate protein-protein interaction in biotechnology and therapeutics. Recently, a designed AR was developed to specifically recognize the surface glycoprotein gp120 as the inhibitor of HIV entry process and virus infection [9]. A stable consensus TPR protein was designed targeting HSP90 with mild affinity [10].

TPR and AR proteins are composed of repeating units of 34 and 33 amino acids, respectively. The basic repeat unit is helix-turn-helix turn in TPR and helix-β turn-helix-loop in AR protein.

Current protein engineering strategies mainly include structure-based rational design and sequence-based design such as directed evolution and consensus design. Consensus design of repeat proteins is focused on the consensus of individual repeats rather than the natural context in creating the templates. It would be useful to understand the structural nature of repeat proteins that define the folding and stability of designed proteins.

In the past two decades, knowledge-based statistical potentials was developed for protein folding and protein structure recognition [11],[12],[13] based on Anfinsen’s thermodynamics hypothesis [14]. Following the concept brought about by Sippl [12],[15], a variety of distance-dependent statistical potentials have been developed [16],[17],[18],[19],[20],[21],[22],[23]. The distance-dependant potential based on Boltzmann equation is given by:

$$\tilde{u}(i, j, r) = -RT\ln\frac{N_{\text{obs}}(i, j, r)}{N_{\text{ref}}(i, j, r)}$$ (1)

Where R is the Boltzmann constant, T is the Kelvin temperature. $N_{\text{obs}}(i, j, r)$ is the observed number of atomic pairs (i, j) within a distance bin r in a database of experimental protein structures. $N_{\text{ref}}(i, j, r)$ is the reference state, which is the expected number of atomic atoms (i, j) in the same distance bin if there is no interaction between atoms.

The main difference of the statistical potentials lies in the selection of reference states. It was suggested that statistical potentials have a contradiction between the universality and pertinence and optimal reference states should be extracted based on specific application environment [24]. Statistical potential represents the pseudoenergy of proteins, therefore can be used to evaluate protein stability.

Unlike globular proteins, the stability of repeat proteins is dominant by the short-range interactions [25],[26]. Multistate kinetic folding pathway studies for some repeat proteins such as TPR and AR proteins disclosed that folding of these proteins is dominated by the competition between the stability of individual repeats and the interactions between repeats [25]. Pluckthun et. al. proposed that folding is a nucleation process, i.e. assembly of a minimal number of repeats triggers the entire folding process [27]. They suggested that the unfolding requires the progressive disruption of the folded repeat and therefore the stability is dependent on the number of repeats. Furthermore, it was suggested that all repeats in repeat proteins are not equal and different repeats have different contribution to stability [25],[28]. Therefore it is necessary to include sufficient features of repeat protein, e.g. distinct repeat proteins with low sequence...
identity and with different protein length, in the statistical potential libraries while calculating the distance-dependant statistical potentials. In order to evaluate the overall stability of repeat proteins and explore their application as novel binding molecules, we developed repeat-specific distance-dependant statistical potential libraries utilizing the structural features of two classes of helical repeat proteins TPR and AR. The structure-based statistical potential opens a way to evaluate the stability of the proteins that are designed by sequence-based approach, and can be used to quickly prioritize the proteins with predicted high stability for subsequent biological function exploitation.

### Materials and Methods

#### All-atom distance-dependant statistical potentials

Distance-dependant statistical potentials are based on the assumption that the three-dimensional structure of a natural protein in its normal physiological environment has the lowest Gibbs free energy [14]. The stability of the proteins was evaluated by the all-atom probability discriminatory function (RAPDF) scoring function [17], which is based on conditional probability function representing preference of atomic distance.

\[
P(C) \times P(d_{ij}^a|C) = P(d_{ij}^a) \times P(C|d_{ij}^a)
\]

where

- \( P(C) \): the probability that any structure picked at random is a member of the “correct” set.
- \( P(d_{ij}^a|C) \): the probability of observing a distance \( d \) between two atoms \( i \) and \( j \) of types \( a \) and \( b \) in a correct structure.
- \( P(d_{ij}^a) \): the probability of observing such a distance in any structure, correct or incorrect.
- \( P(C|\{d_{ij}^a\}) \): the probability the structure is a member of the “correct” set, given it contains the distances \( \{d_{ij}^a\} \).

\( \{d_{ij}^a\} \) is the distance between atoms \( i \) and \( j \), of type \( a \) and \( b \), respectively.

The probabilities of observing the set of distances is expressed as products of the probabilities of observing each individual distance. An approximation is made that all distances are independent of one another, thus

\[
P(\{d_{ij}^a\}/C) = \prod_{ij} P(d_{ij}^a|C) \times P(\{d_{ij}^a\}) = \prod_{ij} P(d_{ij}^a)
\]

From Equations (3) and (4), the following equation can be retrieved:

\[
P(C|\{d_{ij}^a\}) = P(C) \times \prod_{ij} \frac{P(d_{ij}^a|C)}{P(d_{ij}^a)}
\]

Where \( P(C) \) is a constant independent of the conformation for a given amino acid sequence.

Statistical potential is obtained from statistics of experimental protein structures. All the atoms in the proteins are classified as 167 residue-specific heavy atom types [17] and the atomic distances between each atomic pair are calculated. These distances are then assigned to 18
different distance bins with distance cutoff value of 20 Å. Except for the first bin which is 0–3 Å, the bin width of the rest of the bins is set as 1 Å.

The score is given by the following logarithm of conditional probability:

$$S_{ij}^{ab} = - \sum_q \ln \frac{P(d_{ij}^q|C)}{P(d_{ij}^q)} \propto - \ln P(C|\{d_{ij}^q\})$$  \hspace{1cm} (5)

Here

$$P(d_{ij}^q|C) = \frac{N_{obs}(i,j,r)}{N_{obs}(r)}$$  \hspace{1cm} (6)

$$P(d_{ij}^q) = \frac{N_{obs}(i,j)}{N_{total}}$$  \hspace{1cm} (7)

Thus the scoring function becomes:

$$S(\{d_{ij}^q\}) = - \sum_q \ln \frac{N_{obs}(i,j,r)}{N_{obs}(r)}$$  \hspace{1cm} (8)

$$N_{obs}(i,j,r): The number of observed atomic pairs (i, j) of atomic type a and b, within bin r.
N_{obs}(i,j): The number of observed atomic pairs (i, j) of atomic type a and b, within 18 bins.
N_{obs}(r): The number of all observed atomic pairs within bin r.
N_{total}: The number of all observed atomic pairs within 18 bins.

The statistical score of a particular protein is the sum of scores associated with all observed atomic pairs within 18 distance bins.

$$S_{score} = \sum_{ij} s_{ij}^{ab}$$  \hspace{1cm} (9)

Where $s_{ij}^{ab}$ is the statistical potential associated for atomic pairs $(i, j)$ with a value of

$$- \ln \frac{N_{obs}(i,j,r)}{N_{obs}(r)}$$

Database of reference protein structures

Six statistical libraries were constructed using $\alpha$, $\beta$, $\alpha + \beta$ and general proteins, AR proteins and TPR proteins, respectively. The $\alpha$, $\beta$, $\alpha + \beta$ and composite protein structure databases collected from Hobohm’s protein database [29]. The library of $\alpha + \beta$ protein structures was filtered by sequence identity cutoff of 25% and resolution cutoff of 1.5 Å resulting in 1271 proteins. The $\alpha$- and $\beta$- protein collections were filtered by sequence identity cutoff of 25% and resolution cutoff of 3.5 Å, resulting in 1007 $\alpha$- and 288 $\beta$- protein structures. The composite protein database is the sum of $\alpha$, $\beta$- and $\alpha + \beta$ databases. The original RAPDF potential based on a general protein database was also used to evaluate the stability of the proteins [17].

TPR and AR proteins were collected from SCOP [30] and PDB database. These proteins were filtered using sequence identity cutoff of 30% to construct the AR and TPR statistical library statistical libraries, which contain 33 AR proteins and 73 TPR proteins, respectively. PRIDE2 executable [31] was used to determine protein fold similarity and structural
relationship was visualized using Drawtree and Drawgram functionalities in PHYLIP package (version 3.5c) [32]. The Arc of tree in Drawtree was set as 250°.

Construction of decoy protein structures

Different decoy protein structures were collected or prepared to evaluate the efficiency of various statistical potential libraries on differentiation of correct structures from incorrect ones. Misfolded protein structures collected from the Decoy ‘R’ Us website were categorized into α-, β-, mixed α+β proteins and used as decoy structures [33]; for AR and TPR proteins, comparison was made between the natural proteins and their corresponding homology models. Additional comparison was made between designed consensus repeat proteins and their respective template scaffolds.

Homology models were built as decoy set for 8 proteins selected from the AR and TPR protein databases. The selection was made based on the criteria that there is sufficient sequence identity between the query and the template protein and they are evolutionarily relevant species (sequence similarity is between 54% and 86%) (Table 1), thus the native and the decoy proteins have structural relevance. Homology models were built using Modeller (UCSF, USA) and the one with lowest DOPE score was kept for each protein.

Results and Discussion

Statistical potentials based on general and α+β proteins

Recently, we evaluated the stability of self-derived peptides derived from three classes of envelope (E) proteins by two state-of-art statistical scoring functions, dDFIRE and RAPDF[17], [34]. It was found RAPDF based Monte Carlo selection method outperformed dDFIRE method for the beta-sheet Class II E proteins although both scoring functions display similar efficiency for the alpha-helical Class I HIV-1 gp41 and the mixed α+β Class III HSV-1 gB proteins [35]. Therefore in the current research, we developed statistical potential based on RAPDF.

Statistical potential libraries based on α+β proteins as well as a composite database of 2566 proteins that comprises all α, β, and α+β proteins were constructed. 26 proteins and their
misfolded decoy proteins were evaluated using the original RAPDF statistical potential and the potential based on the composite protein database. It was demonstrated that the score difference between the natural structures and their corresponding misfolded decoy structures is similar evaluated by these two general potentials (Fig. 1). Similarly, the stability score difference of 20 α+β proteins and their decoy partners is similar when it is evaluated using the statistical potential based on 1271 α+β protein and the original RAPDF potential (Fig. 2). It is not surprising as the statistical potentials of both the general protein and α+β databases were constructed based on a large dataset of protein structures such that the feature of common proteins was encompassed.

**Statistical potentials based on proteins with certain secondary structure**

**α and β statistical potentials.** Since spatial arrangement of the atoms of proteins is crucial for distance-dependant statistical potential, we propose the feature of certain secondary
structure should be reflected in the specific statistical libraries constructed based on representative protein secondary structures. Statistical potential libraries based on α-, β- proteins were constructed. The stability difference between the natural and misfolded decoy proteins is significantly larger when evaluated by the potentials constructed based on α or β proteins, compared with those evaluated by the general RAPDF potential (Figs. 3 and 4). The stability gap between the natural and incorrect structures is even greater for the dynamic solution structure of the C-terminal domain of cellobiohydrolase I (CT-CBH I), a β protein with two disulfide bonds (pdb code: 1CBH, Fig. 4) [36]. The general statistical potential is inferior to the β potential in identifying the correct conformation from the decoy one indicating the structural feature of the β-protein in particular the disulfide bridges is not sufficiently represented in the general potential library. We also evaluated multiple decoy sets collected from the Decoy ‘R’ Us website [33]. It can be seen that our method is also more effective in discriminating native or near-native from non-native ones (S1 Table).
Therefore it is necessary to use specific statistical potential to evaluate the stability of proteins with certain secondary structure.

**Repeat-specific statistical potentials.** 100 TPR and 68 AR non-redundant proteins were collected from SCOP and PDB database. Using sequence identity cutoff of 30%, 33 AR proteins and 73 TPR and TPR-like proteins were retained to construct the AR- and TPR- specific statistical potentials. Although there are 8,000 AR sequences in the SMART database [37], only 33 AR proteins were identified with less than 30% sequence identity. This is because most of the resolved structures of AR were designed proteins which share high sequence similarity. The number of repeat or repeat-like motifs in the AR or TPR proteins is between 1 and 11.

Pair-wise protein fold similarity comparison was performed for the non-redundant TPR and AR protein database using PRIDE executables and the results were plotted using Drawtree (Fig. 5) and Drawgram (S1 Fig.). We found that the TPR protein library exhibits high diversity with the tree branches spreading around the origin. In contrast, the AR protein library is more populated, with a barren space, where no structure has been deposited. Structural comparison was also performed for the TPR and AR protein libraries filtered by 30% sequence similarity.

![Diagram showing protein stability comparison](image URL)
Interestingly, compared with the TPR library, the proteins in the AR library are generally more similar in structure. Repeat-specific statistical libraries based on two classes of repeat proteins AR and TPR were constructed. Homology models for eight AR and eight TPR proteins were built as decoy structures and the stability difference between the natural proteins and the corresponding homology proteins were calculated using the repeat-specific statistical potentials (Figs. 6 and 7). We selected the templates which share similar sequence identity (54%-86%) to the natural ones to construct homology models as decoys such that they are structurally similar to the natural repeat (correct) proteins.

It was exhibited that the stability difference evaluated by AR or TPR specific statistical potential is remarkably higher than those evaluated by the general, α or β statistical potentials. This indicates the structural feature of the repeat proteins is sufficiently reflected in the statistical potential libraries and the repeat specific statistical potential is efficient in identifying natural repeat proteins from decoy structures even when the difference between the natural and decoy structures is trivial. It is worth noting that the stability difference is undetectable for the (S2 Fig. and S3 Fig.).

Fig 4. Evaluating the stability of β proteins using distance-dependant statistical potential based on β protein library (288 proteins). RAPDF (β) represents the statistical RAPDF scores calculated using the β- database. First 4 sets were single misfold decoy sets and the rest 16 sets were from multiple decoy sets with a representative decoy selected.

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AR domain of Drosophila notch receptor (pdb code: 1OT8: A) [38] and two TPR proteins, the TPR domain of Human Kinesin Light Chain 2 (pdb code: 3CEQ: B) [39] and the TPR palm domain of Menin (pdb code: 3U84: A) [40]. This is because these repeat proteins have high structural similarity to their respective templates (S1 Fig.). In particular, the sequence identity between 3U84 and its template 3RE2 is only around 54% (Table 1), however, their statistical potential scores are indiscernible due to the exceptionally high structural similarity.

Mutation of Arg50 of TPR-containing MamA protein (pdb code: 3AS5) into glutamate (pdb code: 3ASD) resulted in disruption of the salt bridge formed between Arg50 and Asp79 and destabilization of entire TPR1 of the protein [41]. We calculated the stability of the natural and mutant TPR proteins using the TPR-specific potential and found that the natural TPR is more stable than the mutant protein (Fig. 8). This is in agreement with the crystal structure of the R50E mutant, where the electron density for the TPR1 was missing.

Due to the significance of repeat proteins in protein recognition, design of novel repeat proteins as alternative binding molecules to antibodies has become an attractive area in biotechnology. Consensus design is a useful biotechnology approach in constructing novel scaffolds to generate binding proteins with improved binding affinity and specificity.

In design of protein with desired binding activity, it is important to select a template onto which functional residues can be grafted. Consensus design is consensus construction of self-compatible repeat module template, a sequence of most frequent amino acid residues at each position decided by multiple sequence alignment. Two distinct consensus design strategies were used in design of AR and TPR proteins. Consensus AR proteins were constructed by fixing the conserved residues that maintain the repeat structures and randomizing the residues that are involved in target protein interaction [42],[43],[44]. In design of consensus TPR
proteins, the repeat scaffold was modified by introducing functional residues involved in target protein recognition and specific binding.

The TPR-specific potential was used to evaluate the stability of consensus TPR proteins. CTPR3, a designed consensus TPR (pdb code: 1NA0) was reported to be more stable than the template protein phosphatase 5 (PP5) (pdb code: 1P17) [45],[46]. Comparison of the statistical scores of the consensus TPR and the natural TPR manifested that the stability difference is more prominent than rest of the potentials (Fig. 7), in accordance with the experimental observation.

It was reported that the designed AR protein was more thermodynamic stable than the natural structure [42],[43]. The AR-specific potentials were used to evaluate the stability of designed consensus repeat proteins. Compared to the natural AR protein GABPβ1 (pdb code: 1AWC: B) [47], the designed consensus 5-repeat AR protein (E3_5) (pdb code: 1MJ0: A) [43]...
is associated with a lower statistical potential score, indicating it is more stable than the natural one (Fig. 6). The difference of the stability between the consensus and natural AR proteins is most prominent using the AR-specific potential among all the potentials, in accordance with the experimental observation.

Another consensus AR bound with maltose binding protein (MBP) (pdb code: 1SVX: A) is associated with comparable statistical score to that of the natural protein bound with GABPα [44]. Unlike TPR, LRR and WD40s proteins, AR and HEAT were reported to demonstrate
great elasticity when binding with their targets [48], [49]. Thus the stability of AR in the bound complex is probably compromised by the conformational change when it binds to the target. Recently, it was reported the buried surface of protein is responsible for protein-protein binding affinity [50]. The buried surface area of consensus off7 MBP is 611 Å² [44], comparable to that of the natural AR protein in complex with GA binding protein (GABPα) (854 Å²) [47].

Thus the designed AR has similar binding affinity to the natural AR. In our previous study, we suggested that the structural stability of proteins is related to their in situ binding potential to the partner regions [35]. The off7 AR bound with MBP displayed comparable statistical score to that of the natural protein. This provides additional support to our assumption that the binding affinity of proteins is dependent on their stability.

E3_5 [43], E3_19 (pdb code, 2BKG) [51] and NI3C (pdb code: 2QYJ) [52] were designed AR proteins derived from same framework residues. E3_5 and E3_19 have difference
sequences in that residues are different at randomized positions while Ni3C has three full-consensus repeats. Our calculations demonstrated that Ni3C has higher stability compared with E3_5 and E3_19. This is in line with observed high thermostability of Ni3C, attributed to the increased salt-bridge interaction on its protein surface. NMR studies disclosed that unfolding of the C-terminal capping repeat limits the stability of designed ARs [53]. Two mutated forms of Ni3C, Ni3C_Mut5 (pdb code: 2XEE, where the C-terminus was extended by three residues) and Ni3C_Mut6 (pdb code: 2XEH, where three additional charged residues were introduced to Ni3C_Mut5) showed increased stability compared to the originally designed AR protein, attributed to increased buried surface area and additional salt-bridge or H-bond interactions [54]. The initially designed Ni3C is already very stable and the two mutants are slightly more stable than Ni3C. Using the statistical potential developed based on the AR proteins, we found both mutants are associated with higher RAPDF scores. In contrast, none of rest four statistical potentials could differentiate them.

Comparison of statistical scores and equilibrium unfolding free energies

Unlike globular proteins, the stability of repeat proteins is dominated by short-range interactions [25], [26]. Folding kinetics indicated that there is a competition between the intrinsic stability of individual repeats and the interactions between repeats. Designed consensus repeat proteins have identical repeat units and therefore provide an excellent system for investigation of the thermodynamic properties of repeat proteins. Two series of TPR proteins, namely CTPR and CTPRan proteins, which only differ by a double mutation per repeat, were engineered by the Regan and Main groups. The equilibrium unfolding and chemical unfolding of two series of CTPR proteins including seven proteins from the CTPRan series (CTPRa2 to CTPRa10) and two from the CTPR series (Table 2) were investigated. Among them, CTPR2 (pdb code: 1NA3) and CTPR3 have two and three 34-aa identical consensus repeats followed by a solvating helix [47]; CTPRa8 (pdb code: 2AVP) contains eight TPR repeats [26].

We calculated the stability of designed TPR proteins using the statistical potential and correlated the statistical scores with the thermal unfolding. The unfolding was monitored using differential scanning calorimetry (DSC) experiment and the model-independent free energies of unfolding (ΔGD, N) were calculated using the Gibbs-Helmholtz equation [55]. An obvious correlation was observed with a R² value of 0.84 (Fig. 9). Thermodynamic unfolding transition can be described by a 1D homozipper Ising model that treats each arrayed element of a repeat protein as an equivalent independently folding unit with nearest-neighbor pair-wise interactions between those units [26]. The free energies for folding were represented by ΔG0→j (j is the number of α-helices) [56]. We further correlated the statistical scores of CTPRan with ΔG0→j

<table>
<thead>
<tr>
<th>Protein</th>
<th>ΔGD, N [54] (kcal/mol)</th>
<th>ΔG0→j [55] (kcal/mol)</th>
<th>RAPDF (TPR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTPRa2</td>
<td>3.2±0.6</td>
<td>3.3±0.9</td>
<td>-82.66</td>
</tr>
<tr>
<td>CTPRa3</td>
<td>4.8±0.4</td>
<td>6.1±1.2</td>
<td>-127.68</td>
</tr>
<tr>
<td>CTPRa4</td>
<td>4.2±1.1</td>
<td>9±1.5</td>
<td>-171.94</td>
</tr>
<tr>
<td>CTPRa5</td>
<td>6±0.8</td>
<td>11.8±1.9</td>
<td>-217.1</td>
</tr>
<tr>
<td>CTPRa6</td>
<td>7.7±0.2</td>
<td>14.7±2.2</td>
<td>-261.36</td>
</tr>
<tr>
<td>CTPRa8</td>
<td>14.3±1.5</td>
<td>20.4±2.8</td>
<td>-350.81</td>
</tr>
<tr>
<td>CTPRa10</td>
<td>23.4±2.1</td>
<td>26.1±3.5</td>
<td>-605.63</td>
</tr>
<tr>
<td>CTPR2</td>
<td>7.6±1.1</td>
<td>-</td>
<td>-154.01</td>
</tr>
<tr>
<td>CTPR3</td>
<td>12±0.7</td>
<td>-</td>
<td>-163.81</td>
</tr>
</tbody>
</table>

doi:10.1371/journal.pone.0119417.t002
that was calculated from fitting into the Ising model. A very strong correlation efficient $R^2$ of 0.93 was also observed. This is reasonable since the free energy is strongly correlated with the number of repeat units [25]. Whereas no correlation was found between the statistical scores and the unfolding energies for general globular proteins (S2 Table). The high correlation...
between the statistical scores and the equilibrium thermal/chemical unfolding free energies of repeat proteins suggests the statistical potential developed here can be accurately used to predict the stability of designed repeat proteins along the multistate kinetic folding pathways.

In consensus design or directed evolution, proteins are engineered so as to have admirable functions such as binding specificity or thermal stability. The designed libraries are usually large with the designed proteins being similar to the original scaffold. The statistical potential developed here can be used to quickly prioritize proteins in the libraries for subsequent functional assessment.

Conclusions

Our research demonstrated that distance-dependant statistical potential is sensitive to the secondary structures. It is necessary to use the specific statistical potential based on specific protein secondary structure database to discriminate between correct and incorrect three-dimensional structures for a given sequence. We demonstrated that the repeat-specific statistical potentials we developed are efficient in differentiating the correct repeat protein structures from incorrect models. The statistical score correlate perfectly with equilibrium thermal/chemical unfolding free energy, and therefore would serve as a novel tool in quickly prioritizing designed repeat proteins with high stability.

The feature of repeat proteins allows for the evolution in biotechnology not only by mutation, but also by inserting, deleting, or shuffling the repeat motif, resulting in large combinatorial libraries. The repeat-specific distance-dependant statistical potentials can be used to rank stability of designed repeat proteins thus would provide guidance to prioritize repeat proteins from the designed combinatorial libraries based on their stability, in order to further explore their potential function in mediating protein-protein interactions.

A web server ‘Stability of Repeat Proteins’ (StaRProtein) is freely accessible via the URL http://StaRProtein.ch.qub.ac.uk. StaRProtein server is an on-line platform for evaluating protein stability, which is based on all-atom distance-dependant statistical potentials. Proteins with different secondary structures including alpha-, beta-, alpha-beta- and repeat proteins such as ankyrin repeat (AR) proteins and tetratricopeptide repeat (TPR) proteins are assessed using specific statistical potentials. Users can upload a protein structure in pdb format and designate the type of statistical potential library file. A statistical score which indicates the stability of the protein, the statistical potential library used and the length of the protein will be returned in output.

Supporting Information

S1 Fig. PRIDE2 structure comparison of non-redundant repeat proteins (Drawgram). The repeat proteins are divided into branches, which are shown as groups (A) AR (B) TPR. (PDF)

S2 Fig. PRIDE2 structure comparison of repeat proteins with less than 30% sequence identity (Drawtree). The repeat proteins are divided into branches, which are shown as groups (A) AR (B) TPR. (PDF)

S3 Fig. PRIDE2 structure comparison of repeat proteins with less than 30% sequence identity (Drawgram). The repeat proteins are divided into branches, which are shown as groups (A) AR (B) TPR. (PDF)
S1 Table. Statistical scores of multiple decoy proteins with different second structures. (PDF)

S2 Table. Comparison of kinetic energies and RAPDF scores of globular proteins. (PDF)

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Author Contributions

Conceived and designed the experiments: MH. Performed the experiments: YX XZ MH. Analyzed the data: MH YX XZ. Contributed reagents/materials/analysis tools: YX MH. Wrote the paper: MH.

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