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A General Computational Approach for Repeat Protein Design

Fabio Parmeggiani^{1,†}, Po-Ssu Huang^{1,†}, Sergey Vorobiev², Rong Xiao^{3,4}, Keunwan Park¹, Silvia Caprari¹, Min Su², Seetharaman Jayaraman², Lei Mao^{3,4}, Haleema Janjua^{3,4}, Gaetano T. Montelione^{3,4}, John Hunt², and David Baker^{1,5,6}

¹Department of Biochemistry, University of Washington, Seattle, WA 98195, USA

²Department of Biological Sciences, Northeast Structural Genomics Consortium, Columbia University, New York, NY 10027, USA

³Center for Advanced Biotechnology and Medicine, Department of Molecular Biology and Biochemistry and Department of Biochemistry, Robert Wood Johnson Medical School, Rutgers, The State University of New Jersey, Piscataway, NJ 08854, USA

⁴Northeast Structural Genomics Consortium, Rutgers, The State University of New Jersey, Piscataway, NJ 08854, USA

⁵Institute for Protein Design, University of Washington, Seattle, WA 98195, USA

⁶Howard Hughes Medical Institute, University of Washington, Seattle, WA 98195, USA

Abstract

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Central

Repeat proteins have considerable potential for use as modular binding reagents or biomaterials in biomedical and nanotechnology applications. Here we describe a general computational method for building idealized repeats that integrates available family sequences and structural information with Rosetta *de novo* protein design calculations. Idealized designs from six different repeat families were generated and experimentally characterized; 80% of the proteins were expressed and soluble and more than 40% were folded and monomeric with high thermal stability. Crystal structures determined for members of three families are within 1 Å root-mean-square deviation to the design models. The method provides a general approach for fast and reliable generation of stable modular repeat protein scaffolds.

Keywords

repeat proteins; computational design; thermodynamic stability; idealized proteins; de novo design

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Correspondence to David Baker: Department of Biochemistry, University of Washington, Seattle, WA 98195, USA. [†]F.P. and P.-S.H. contributed equally to this work.

Present address: S. Caprari, Max Planck Institute for Informatics, 66123 Saarbrücken, Germany.

Accession numbers

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Introduction

Repeat proteins play key roles in biological processes ranging from adhesion to signaling to defense mechanisms [1]. These proteins consist of adjacent series of usually non-identical repeated amino acid sequences; in most cases, these repeated units fold cooperatively into either a solenoid-shaped or a toroid-shaped structure [2–4]. Although extremely diverse in structure and sequence, repeat proteins are characterized by short-ranged intra-repeat and inter-repeat interactions between residues proximal in sequence [2]. The intrinsic modularity of repeat proteins allows combination of functionalities in a single domain (e.g., recognition motifs for nucleic acids [5] and peptides [6]) and can be used to generate biomaterials with tunable mechanical properties [7]. However, interactions between neighboring repeats are not always conserved; hence arbitrary extension by repeat insertion is not usually possible.

To allow modular extension, we have generated designed repeat proteins with selfcompatible repeating elements using consensus-based approaches [8–18]. Consensus sequences are defined by the most common residue at each position in a multiple sequence alignment (MSA) of the proteins or repeats in a family. This approach is conceptually simple and powerful but does have non-optimal features. First, the consensus sequence can vary depending on the collection size and the selection method for the sequences used in the alignment. Second, residue–residue packing, particularly critical in the formation of a uniquely defined hydrophobic core, is not considered, and hence in some cases, the consensus may have sub-optimal residue–residue interactions. Incorporating amino acid covariation information derived from statistical analysis of naturally occurring sequences can capture some of these residue–residue coupling effects [19–21], but reliable estimates of covariance require large numbers of sequences that are not available for all protein families.

Here we describe a general computational approach for repeat protein design that integrates Rosetta *de novo* structure generation and design methodology with protein family-based sequence and structural information. By automatically generating very low energy design models compatible with the available sequence and structure information, the method provides increased versatility compared to standard sequence consensus-based approaches and reduces the manual intervention required to achieve stable designs..5

Results and Discussion

We developed a computational approach that integrates sequence and structural information with Rosetta [22] *de novo* folding and design calculations for the generation of idealized repeat proteins (Fig. 1). Families with α helical, β and mixed α/β secondary structure were chosen for redesign to illustrate the generality of the method. Sets of sequences were designed for six protein families: ankyrin, armadillo, tetratrico peptide repeat (TPR), HEAT, leucine-rich repeats (LRR) and WD40.

Overview of computational method

For a repeat protein family of interest, protein structure and sequence information are extracted from publicly available databases and implemented as constraints in the Rosetta modeling suite [22]. Residue–residue distance constraints are used to guide Rosetta structure

generation calculations. Sequence constraints derived from MSAs are used to bias Rosetta design calculations (see Materials and Methods).

Protein backbones are generated *de novo* [23–25] as poly-valine to avoid bias toward a particular template structure. Structure-based and sequence-based constraints guide the sampling and limit the search space to an area defined by the protein family. We exclude from the sequence and structural alignments sequences that were previously designed with consensus-based approaches. Proteins are represented as a single chain formed by a series of identical repeats. The number of repeats and the secondary structure can be arbitrarily chosen or derived from existing proteins.

Backbone conformations are generated by Monte Carlo fragment insertion using RosettaRemodel [25], with insertions in a single repeat replicated in all other repeats (Fig. 1a). The trajectories optimize a function consisting of the Rosetta energy supplemented with family-specific structural constraints (Fig. 1b). Rosetta sequence design calculations are then carried out on the low-energy backbones that satisfy the constraints. The sequence is designed simultaneously in all the repeats (Fig. 1c), using the Rosetta energy function with a sequence profile term favoring residues observed in the family MSA (Fig. 1d).

Most native repeat proteins form a solenoid-like structure with specialized terminal repeats to avoid aggregation and low solubility. Exposed hydrophobic residues in the terminal repeats were substituted with polar amino acids using Rosetta design calculations (Materials and Methods).

Sequence logos [26,27] of the designed proteins for each family are compared to their native counterparts in Fig. 2. The most conserved positions were recapitulated with a few exceptions in the HEAT family (see below in the family description). Even in the absence of MSA constraints, structural constraints alone allowed recovery of the key residues for each fold (Supplementary Figs. S1 and S2 and Table S1). For experimental evaluation, the full protocol with both sequence and structural constraints was used to generate the final sequences.

Synthetic genes were constructed for several low-energy designs from each protein family (Table 1), and the designed proteins were expressed in *Escherichia coli*. Proteins were purified from the soluble fraction of cell lysate and their oligomeric state was characterized by analytical gel filtration (AGF) with multi-angle light scattering (MALS). Secondary structure and cooperative unfolding during thermal denaturation were measured by circular dichroism (CD) (Table 2 and Fig. 3). We solved crystal structures of several of the designs and their agreement with the models is shown in Fig. 4.

In the following sections, we describe the computational and experimental results for each family.

Ankyrin (ank)—Ankyrin repeat proteins are characterized by a short β hairpin and two antiparallel α helices (Fig. 3). The design calculations converged on set of sequences with about 50% of the residues conserved (see sequence logo for designed ankyrin; Fig. 2). All six ankyrins selected for experimental characterization were expressed solubly with high

yield (>50 mg/l of culture), monomeric, α helical by CD and stable up to 95 °C (Fig. 3). The same properties were observed in previously designed ankyrin repeats (DARPins) [8]. The accuracy of the designs was confirmed by X-ray crystallography. The computational models were within 1 Å root-mean-square deviation (RMSD) from their corresponding crystal structures (Fig. 4a).

Armadillo repeat proteins (arm)—The basic repeat unit of this family is formed by three helices located roughly in the same plane. The designs recapitulated all the sequence features of the family and converged to specific amino acids in several positions (Fig. 2). Surface residues were optimized to reduce the potential electrostatic repulsion from repeated charged amino acids, without affecting significantly the overall energy. Seven out of eight designs tested were expressed and soluble and three were found to be monomeric and folded (Table 2 and Fig. 3). The crystal structure of design arm8 was determined and proved to be very close to the design model (1 Å RMSD; Fig. 4b).

Tetratrico peptide repeats (TPR)—The TPR repeat unit is formed by two α helices connected by a short loop. The design calculations did not converge into a narrow sequence space as in the ankyrin case but did capture the key features of the family (Fig. 2). Capping repeats were not modified, as there were no large exposed hydrophobic residues in the designs. Six 4-repeat designs were selected for experimental testing. All the proteins were expressed at a yield above 30 mg per liter of culture. One displayed molten globule-like properties and five were folded. Four of these were monomeric, while the fifth formed a dimer in solution. TPR3 displayed a cooperative and reversible thermal unfolding, while the other four folded proteins did not denature at 95 °C (Fig. 3).

HEAT repeat proteins (HEAT)—HEAT proteins form a large family of α solenoids characterized by two main helices connected by a loop. The remainder of the repeat can assume multiple conformations, from straight helices to kinked helices to long loops [28]. This variability represents a challenge for consensus-based methods and only one successful design from a small and very conserved subgroup has been reported so far [12]. The designed sequences recapitulate the conserved sequence features with the exception of Pro9 and Arg23; Asp17 was only partially recovered (Fig. 2). The structural features associated with these residues are poorly sampled and energetically disfavored during *de novo* backbone generation. Instead, the design protocol explored alternative solutions to lower the total energy and improved packing by substituting Pro9 within the first α helix with residues with higher helix propensity and by replacing a buried salt bridge formed by Asp17 and Arg23 with alternative packing solutions. Eight designs were experimentally tested. Three designs were soluble and had the expected CD spectrum, but only HEAT7 was expressed at high yield and monomeric (Table 2 and Fig. 3).

Leucine-rich repeat proteins (LRR)—LRRs are characterized by repeats containing a β strand packed on an α helix (or 3–10 helix or loop) that form a horseshoe-shaped $\alpha\beta$ solenoid. The length of the repeat can vary, with 24 residues the most common size. Two short β strands extracted from variable lymphocyte receptor (VLR) structure 206S (residues 103–105 and 127–128, chain A) were used to ensure the correct pairing during backbone

design. Sequence refinement led to the sequence family depicted in Fig. 2, with all the conserved residues recovered. The two lowest-energy models were similar to natural and designed VLRs [13]. Surface residues were optimized without MSA bias to reduce the electrostatic repulsion from repeated charged amino acids. We used the N-terminal capping repeat of internalin B, successfully employed in VLRs [13], while the C-terminal cap was generated from the internal repeat unit. The two experimentally characterized LRR designs were stable and folded, although a large fraction formed soluble aggregates in addition to monomers (Fig. 3). In contrast to other designs, they aggregated irreversibly at high temperature. We solved crystal structures of two of the designs. These structures obtained are very similar to the design models (~1.1 Å RMSD across backbone heavy atoms in the internal repeats) (Fig. 4c). Although the designs share the same N-terminal cap and are more than 80% identical in the internal repeat assumes an alternative conformation, forming an α helix that packs against the exposed core of the last internal repeat.

WD40

WD40 proteins form a toroid β propeller, where each "blade" is a four-stranded β sheet that packs against the two neighboring sheets. The fold is characterized by buried polar interactions that were disfavored in the initial design calculations. The definition of the WD40 repeat in the SMART and Pfam databases does not correspond to the structural unit but instead comprises three strands and the fourth strand of the previous blade [29,30]. However, a repeat definition matching the structural unit combined with explicit buried polar residues (Trp24, Asp25 and Ser/Thr14) led to a pool of sequences resembling closely the native proteins (Fig. 2). Separate design calculations for 6-bladed, 7-bladed and 8-bladed propellers were carried out. Thirteen designs were experimentally tested and nine were found to be expressed and soluble; however, only one with eight repeats, WD40 847 appeared folded (Table 2 and Fig. 3). WD40 847 was purified as large soluble aggregates, but upon heating for 10 min at 80 °C, it became mainly dimeric. The CD spectrum was closely resembled by the spectra observed for existing β propellers [31]. WD40_847 was expressed also as a half-propeller, with only four repeats. AGF-MALS analysis identified the species in solution as dimer, suggesting that it might indeed form an 8-repeat β propeller, but we were unable to confirm this with an X-ray crystal structure.

Contribution of protein family information to designed sequences

To investigate the contributions of the different information sources to the design models, we carried out calculations in which either the structural constraints or the sequence constraints were eliminated. At the backbone building stage, structural constraints significantly increase the fraction of models with structural similarity to the naturally occurring family members (Materials and Methods and Supplementary Fig. S1). For each repeat protein family, a subset of the models satisfies the topology requirements, even in absence of structural constraints. The gain is family dependent, with only marginal improvements for simple two-helix topologies such as TPR to more than 90% for a more complex three-helix topology such as armadillo.

The effect of the MSA-derived sequence constraints on the designs generated by our protocol was evaluated by comparing the sequence profiles of models obtained with or without sequence constraints (Materials and Methods). Profile–profile comparison to Pfam families using HHsuite [32] shows that the constraints increase the similarity of the profile to the original family from 35% to 80%, on average (Supplementary Fig. S2). With the exception of the HEAT family, where alternative combinations of core residues were explored, even in the absence of sequence information, the closest match to the designed protein sequence profile in the Pfam database is the naturally occurring corresponding repeat protein family (Supplementary Table S1).

Conclusions

The approach presented here generalizes the current MSA-based methods for repeat protein design with an automated pipeline that integrates sequence, structure and energetic information. Designing backbones *de novo* avoids potential bias due to the use of a single or few template structures.

Forty percent of the proteins designed with our method were folded and had a melting temperature ($T_{\rm m}$) of 57 °C or greater (Table 2). The crystal structures we were able to solve had an RMSD of about 1 Å to the design models. Rosetta calculations recapitulate the majority of the sequence features of all the six families and generate models with excellent core packing and backbone geometry. Family-specific structures, such as proline kinks and conserved buried hydrogen bond networks or charge interactions, can be enforced using specific local conformation of the backbone or a set of residue identities.

The use of general sequence and structural constraints allows greater exploration of the sequence space available for repeat protein families than strict consensus-based approaches. As shown in Fig. 5, the method generates low-energy sequences that differ from the consensus and were not explored in previous successful designs, in particular, for TPR and armadillo families [10,11], where variations also occur in several hydrophobic core residues. Although the selected amino acid at each position is usually among the three most frequent, the discrimination of viable sequences among the vast number of potential combinations represents a challenge for simple consensus-based approaches. In the armadillo case, the first consensus design (armC in Fig. 6) had molten globule-like characteristics and stabilization required a second round of refinement (armM in Fig. 6) [10]. In contrast, three of our designed sequences (arm1, arm3 and arm8 in Fig. 6) with only ~60% sequence identity to the consensus are stable up to 95 $^{\circ}$ C, compared to a $T_{\rm m}$ of 70 $^{\circ}$ C observed for armM. Overall, our protocol expands the traditional consensus-based approaches, reproducing the finding for highly conserved families, and offers a general solution to the design of idealized repeat proteins, producing a broader range of sequences than what would be available through consensus design.

The extension of Rosetta *de novo* design methods [23,24] to repeat protein architectures allows the computational design of extended modular non-globular structures for a wide range of applications. In the limit of no available data for a particular topology, it should be

possible to use this approach to design completely new types of repeat protein structures and sequences.

Materials and Methods

Generation of sequence and structural constraints

The repeat consensus sequences were obtained from family alignments in the SMART database [29,33]. For the HEAT family, not present in SMART, the Pfam [30] seed alignment was used. A double-repeat sequence was generated by duplication of the consensus. When the consensus sequence did not cover the whole repeat, connecting fragments were added using alanine residues as placeholders. The length of this linking sequence was based on the shortest connection observed, with at least a 10% frequency, in repeats from crystal structures of family members. Using this sequence, we obtained an improved double-repeat consensus and a sequence profile after five rounds of PSI-BLAST [34,35]. Sequences previously designed were excluded from the database. The PSI-BLAST profile was implemented in Rosetta as Sequence-Profile constraints. According to Eq. (1) below, at each position, the amino acid frequencies (*f*, ranging from 0 to 100) were converted to arbitrary Rosetta energy units (REU) using pseudocounts to allow cases when particular amino acids were not observed.

$$REU = -\log\left((f_i+1)/\sum_i (f_i+1)\right) \quad (1)$$

The values are positive, representing an energy penalty, with higher values for less frequent amino acids, with a maximum value of 2.08 REU.

We selected 100 template structures from the PDB repository[‡] (from Refs. [36] and [37]) using HHsearch [38]. Previously designed structures were excluded from the template list. The repeat consensus was threaded on the templates and the resulting models were clustered with 2 Å RMSD cluster radius using Rosetta [22]. For each model within the largest cluster, distances of carbon α to carbon α (C^{α}–C^{α}) between residues were measured. Contacts were defined as distances shorter or equal to 10 Å (distance cutoff) between amino acids at least 4 residues apart (sequence separation). C^{α}–C^{α} contacts present in all models were considered as conserved and the average distance was calculated. The conserved intra-repeat and interrepeat contacts were implemented in Rosetta as AtomPair constraints, described by harmonic functions centered at the average C^{α}–C^{α} distances, with a spring constant of 10. For each double-repeat model, about 190 AtomPair constraints were used on average, including intra-repeat and inter-repeat constraints.

Secondary structures were assigned as helix, strand or loop, using dssp [39], to the templates belonging to the selected cluster employed for constrained generation. The most common secondary structure at each position was chosen and the information was implemented as Rosetta blueprint file.

[‡]www.wwpdb.org and www.rcsb.org.

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Backbone design and refinement

Protein backbones were generated by insertion of fragments of 3 and 9 residues using RosettaRemodel [25]. A definition of the repeat secondary structure derived from the existing protein was used as starting input. The chain was represented as poly-valine during this initial stage, followed then by sequence design. Generation of repeat proteins required the implementation of simultaneous insertion of fragments in each repeat, as well as coupling of dihedral angles and side chains identities between repeats. The structure-based and sequence-based restraints were used to guide the search space toward the desired fold. About 5000 backbone models per family were generated and clustered by RMSD using the cluster application in Rosetta [22] with a cutoff of 3 Å. The quality of the models was evaluated in comparison to the average conserved interatomic distances within proteins in the family, expressed as AtomPair constraints as described above, using a harmonic function with flat bottom between average \pm standard deviation and spring constant of 10. Violations of these constraints were calculated as energetic penalties. Structures with violations up to 5 REU per repeat, corresponding to less than 10% of the total energy of a correct full atom model or crystal structure, were accepted and their frequency is depicted in Supplementary Fig. S1.

Protocols described as Rosetta_scripts [40] were used for refining the sequences. Three cycles of sequence design and backbone minimization were performed while optimizing the packing interactions and the total energy. The available amino acids for each position were restricted based on the secondary structure and the solvent accessibility [24,25]. Cysteines were excluded to prevent formation of oligomers upon oxidation. The lowest-energy models from the 30 most populated clusters were selected for refinement, with 500 trajectories each.

Designs were filtered according to total energy (designs within 15% of lowest-energy model), and values of chi2 dihedral angles of aromatic residues ($70^{\circ} < chi2_dh < 110^{\circ}$) [24]. Final designs with lowest energy and RosettaHoles score [41] of <0 were selected. RosettaHoles values up to 1 were accepted for WD40 to increase the number of potential candidates. When several designs with similar values were available, sequence composition (e.g., low number of alanines in the core) was used as discrimination factor.

Hidden Markov model profile–profile comparison was carried out using HHsuite [32] with default settings and the Pfam database [30] (Supplementary Table T1). To visualize the influence of sequence constraints on the selected sequence pool, we normalized the score values by the maximum scores obtained by self-alignment of the reference Pfam family, according to Kamisetty *et al.* [42] (Supplementary Fig. S2).

Design of capping repeats

Solenoid-like repeat proteins are often characterized by N-terminal and C-terminal specialized capping repeats that protect the hydrophobic core from solvent exposure. The low sequence conservation and the variability in conformation of capping repeats within families prevented the use of a reliable sequence profile as a general strategy for design of capping repeats; hence the Rosetta energy function alone was used to guide selection of surface residue identities. Capping repeats were designed from the internal repeat by

mutating exposed hydrophobic residues into polar residues, except for leucine-rich repeat (LRR) N-terminal cap where internalin B (residues 25–110) was used as in VLR designs [13]. The first internal repeat was modified to be compatible with the grafted cap. The backbones of the final models within ankyrin and armadillo families were all very similar. The most frequent amino acids replacing the exposed hydrophobic residues in the simulations were used in all models of the family to generate the capping repeats.

As observed in native sequences and in previous designs, the N-terminal ankyrin capping repeat contains 3 helical N-terminal residues [8,43], and the N-terminal armadillo repeat contains only two helices [10]; therefore, these capping repeats were modifying accordingly. TPR repeats did not possess large exposed hydrophobic residue; therefore, no specialized capping repeats were introduced.

Surface refinement and manual intervention

LRR and armadillo final sequences were characterized by a few charged residues in close proximity. A final design round of the surface residues was performed without sequence constraints, but the positions were selected following structure examination, instead of automatically. For LRRs, positions 1, 3 and 19 were redesigned, while one or two selected positions per repeat (26, 30 or 34) were changed in armadillo sequences from glutamate, lysine or arginine to glutamine in 6 out of 8 models.

Molecular biology and biochemistry

Gene synthesis and cloning—Genes were synthesized and cloned in vector pET21_NESG by GenScript (Piscataway, NJ). gBlocks for TPR genes and oligonucleotides were synthesized by Integrated DNA Technologies, Inc. (Coralville, IA) and cloned into pET21_NESG. WD40 genes were synthesized by gen9 (Cambridge, MA) and cloned into pet15_NESG vector via Gibson cloning [44]. Cloning strains used were XL1-blue and XL10-gold (Agilent Technologies).

Protein expression and purification—Proteins were expressed in BL21(DE3) *E. coli* cells (Life Technologies) at 37 °C and induced with 250 μM isopropyl-β,D-thiogalactopyranoside (IPTG), either overnight at 22 °C or for 4 h at 37 °C, without significant difference in yield. For cases with low growth or lack of expression, BL21(DE3) pLysS (Life Technologies) was used, without any significant improvement. Cells were lysed by sonication and the clarified lysate was loaded on a Ni-NTA superflow column (Qiagen). Lysis and washing buffer was 50 mM Tris (pH 8), 500 mM NaCl, 30 mM imidazole and 5% (v/v) glycerol. Lysozyme (2 mg/ml), DNase I (0.2 mg/ml) and protease inhibitor cocktail (Roche) were added to the lysis buffer before sonication. Proteins were eluted in 50 mM Tris (pH 8), 500 mM NaCl, 250 mM imidazole and 5% (v/v) glycerol and were dialyzed overnight in 20 mM Tris (pH 8), 500 mM NaCl or PBS (12 mM phosphate, 137 mM NaCl, 2.7 mM KCl, pH 7.4). Protein concentrations were determined using a NanoDrop spectrophotometer (Thermo Scientific). Except indicated above, enzymes and chemical were purchased from Sigma-Aldrich.

Biophysical characterization—Secondary structure content and thermal stability were monitored by CD using an AVIV 62S DA spectrometer (Aviv Biomedical, Lakewood, NJ). Thermal denaturation was followed at 220 nm for structures containing α helices, at 212 nm for LRR and at 215 nm for WD40. Oligomeric state was assessed by AGF coupled to MALS. A Superdex 75 10/300 GL column (GE Healthcare) equilibrated in PBS was used on a HPLC LC 1200 Series (Agilent Technologies) connected to a miniDAWN TREOS (Wyatt Technologies). The chromatograms shown in Fig. 3 were recorded for 100 µl samples at 1–4 mg/ml concentration, with a flow rate of 0.5 ml/min. Protein molecular weights were confirmed by mass spectrometry on a LCQ Fleet Ion Trap Mass Spectrometer (Thermo Scientific).

Preparation of protein samples for crystallography—Crystallization was attempted for all the folded designed repeat proteins. The plasmids were transformed into *E. coli* BL21(DE3) pMgK-competent cells. All proteins were expressed and purified based on the standard procedures of Northeast Structural Genomics to produce selenomethionine-labeled samples for X-ray crystallography [45]. The selenomethionine-labeled proteins were grown at 37 °C in MJ9 minimal media [46]. When the OD₆₀₀ reached 0.6, selenomethionine, lysine, phenylalanine, threonine, isoleucine, leucine and valine were added 10 min before induction with 1.0 mM IPTG [47]. Protein expression was carried out at 17 °C. Following overnight incubation, we harvested the cells by centrifugation and stored at –80 °C.

Cells were resuspended in 30 ml of lysis buffer [50 mM Tris (pH 7.5), 500 mM NaCl, 40 mM imidazole, 1 mM TCEP and 0.02% (w/v) NaN₃]. Following lysis by sonication, we loaded the supernatant onto an ÄKTAxpress system (GE Health-care) using a two-step protocol consisting of ion metal affinity chromatography (HisTrap HP, 5 ml) chromatography followed by gel-filtration (HiLoad 26/60 Superdex 75) chromatography. Protein-containing fractions were pooled and concentrated to a range of 7.8–12.35 mg/ml. Protein purity and molecular mass were evaluated using SDS-PAGE and matrix-assisted laser desorption/ionization/time of flight mass spectrometry. These pET expression vectors (OR264-21.1, OR265-21.1, OR266-21.1, OR267-21.1, OR329-21.1, OR464-15.1 and OR465-15.1) have been deposited in the PSI Materials Repository[§].

Samples for crystallization were assessed by AGF with static light-scattering detection (AGF-MALS). Protein samples at 20 mM Tris–HCl (pH 7.0), 100 mM NaCl, 5 mM DTT and 0.02% NaN₃ were injected onto an analytical gel-filtration column (Shodex KW-802.5; Shodex, Japan) at room temperature. The HPLC was run on an Agilent series 1200 system at a flow rate of 0.5 ml/min. Data were then collected on a miniDAWN (TREOS) light-scattering instrument (Wyatt Technology) and refractive index (Optilab rEX). The data were analyzed with ASTRA software (Wyatt Technology, version 6.1.1.17).

Structure solution and refinement—Crystallization screening was performed using a microbatch-under-oil crystallization method at 4 °C (OR265, OR267 and OR329) or 18 °C (OR264, OR266, OR464 and OR465) [48]. After optimization, protein crystals useful for structure determination were grown in drops composed of 1.0 µl of protein and 1.0 µl of

[§]http://psimr.asu.edu/.

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precipitant solution [40% polyethylene glycol (PEG) 1000, 0.1 M lithium chloride and 0.1 M Taps (3-{[2-hydroxy-1,1-bis(hydroxy-methyl]amino}-1-propanesulfonic acid), pH 9.0 (OR264); 40% PEG 400, 0.1 M lithium sulfate and 0.1 M Hepes, pH 7.5 (OR265); 40% PEG 1000, 0.1 M potassium nitrate and 0.1 M Hepes, pH 7.5 (OR266); 20% PEG 4000, 0.1 M magnesium sulfate and 0.1 M Tris-HCl, pH 8.0 (OR267); 20% PEG 8000, 0.1 M magnesium nitrate and 0.1 M sodium citrate, pH 4.2 (OR329); 25% PEG 3350 and 0.1 M Bistris (2-[bis(2-hydroxyethyl)amino]-2-(hydroxy-methyl)propane-1,3-diol), pH 5.5 (OR464); 40% PEG 4000, 0.1 M potassium phosphate monobasic and 0.1 M Mes (4morpholineethanesulfonic acid), pH 6.0 (OR465)] under paraffin oil (Hampton Research). Data sets were collected at beamline X4A (OR265, OR266, OR329 and OR464) or X4C (OR264, OR267 and OR465) at the National Synchrotron Light Source at 100 K. The diffraction data from single crystals were processed with the HKL2000 package [49]. The structures were solved by molecular replacement using program MolRep [50] and models 2XEE (OR266) and 4GPM (OR267) or using BALBES [51] and models 2XEE (OR264), 4HB5 (OR265), 4DB8 (OR329) and 3RFJ (OR464 and OR465). The models were completed using iterative cycles of manual rebuilding in Coot [52] and were refined with the program PHENIX [53]. The quality of the model was inspected by the program PROCHECK [54]. The data processing and refinement statistics are provided as Supplementary Materials.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations used

MSAmultiple sequence alignmentAGFanalytical gel filtrationMALSmulti-angle light scatteringVLRvariable lymphocyte receptorPEGpolyethylene glycol

Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.jmb. 2014.11.005.

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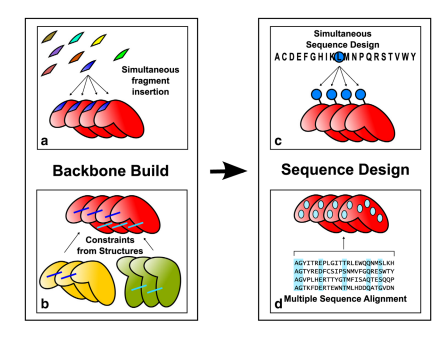


Fig. 1.

Overview of repeat protein design protocol. The protein backbone is built by simultaneous fragment insertion (a) guided by constraints derived from existing structures (b). Rosetta sequence design calculation is carried out enforcing sequence identity between repeats (c) and amino acids at specific positions are favored according to their frequency in the MSA of the target family (d).

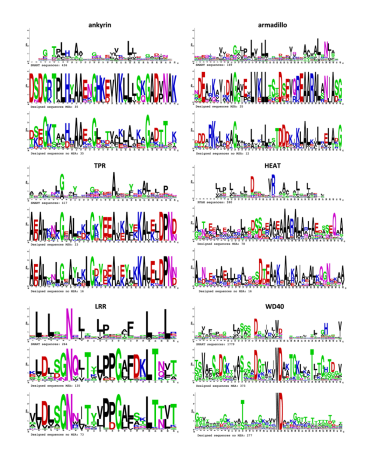


Fig. 2.

Comparison of computationally designed and existing sequences. The sequence logo of naturally occurring sequences (top logo for each family) is compared to the sequence logo obtained from the pool of computationally designed sequences with (middle) or without (bottom) family-specific MSA. Blank entries in the logo of naturally occurring sequences are positions not covered by the MSA available in the database. WD40 designs included in both cases an additional sequence bias at positions 14, 24 and 25 (see the family description in the results section), which accounted for the high recovery in the profiles. As noted in the text, the WD40 sequences were rearranged to match the structural unit in the protein.

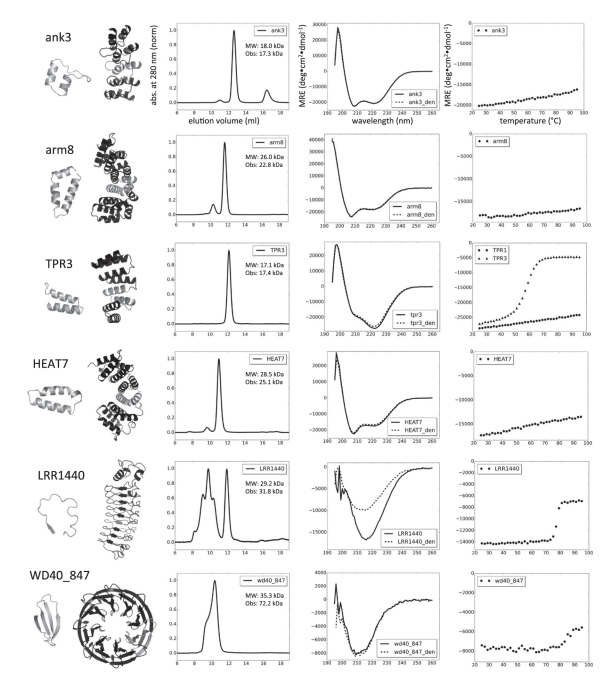


Fig. 3.

Characterization of designed repeat proteins. From left to right, repeating unit and protein model, AGF, CD spectrum and thermal denaturation profile. The repeating unit is highlighted in gray in the models. Axis labels in the first row apply also to the other plots. Abs indicates normalized absorbance measured at 280 nm; MRE is mean residue ellipticity. AGF was performed on a Superdex 75 column (void volume at 8 ml); MW is the expected molecular weight and Obs indicates the observed mass in MALS for the main non-aggregating peak. CD spectra were recorded at 25 °C before and after denaturation (_den). All proteins were able to refold upon thermal denaturation with the exception of LRRs.

Thermal denaturation was followed at 220 nm for ank, arm, TPR and HEAT; thermal denaturation was followed at 212 nm for LRR and at 215 nm for WD40. Data shown are from one representative monomeric protein for each family. An additional temperature denaturation curve is displayed for TPR, showing the different behavior observed among monomeric designs within the family.

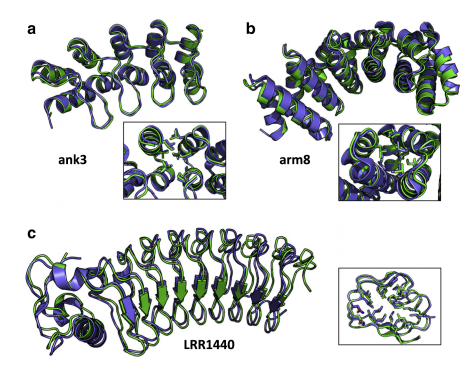


Fig. 4.

Superposition of models and crystal structures for ank3 (a) (RMSD of 0.9 Å), arm8 (b) (RMSD of 0.9 Å) and LRR_1440 (c) (RMSD of 1.1 Å). Models are in green and crystal structures are in blue. In most cases, the core residues assume the conformation predicted in the models, as shown in (a), (b) and (c) insets for some of the side chains. Parts of the structures have been removed to display the core residues. RMSD was calculated using backbone heavy atoms. For LRR, the N-terminal capping repeat was not included in the RMSD calculation, when considered that the RMSD increases to 1.6 Å. Pictures were realized with PyMOL (Schroedinger).

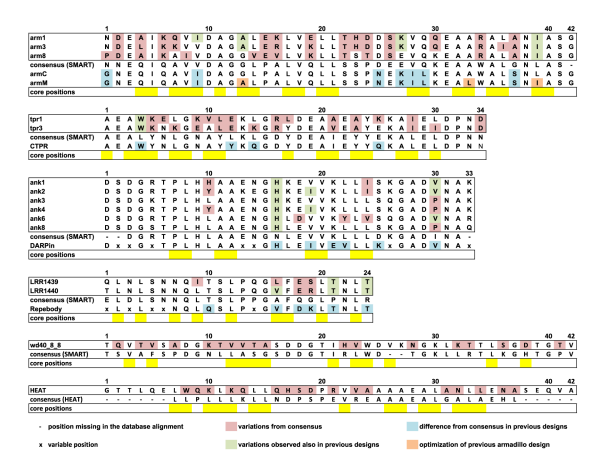


Fig. 5.

Comparison of designed repeat protein sequences to strict consensus sequences and previous consensus-based designs. Armadillo, TPR, ankyrin and LRR designed sequences are compared to consensus sequences from SMART database [29,33] and to previous successful designs armC and armM [10], CTPR [11], DARPin [8] and Repebody [13]. HEAT and WD40 designed sequences are compared to the consensus from Pfam and SMART, respectively. Positions missing in the SMART and Pfam alignments are represented as dashes and were not considered. Positions identified as variable and included in library designs are labeled with *x* and were also not considered. Pink: positions that vary between the designs presented here and the SMART consensus; green: positions that differ from the consensus but present in previous designs; blue: differences between the previous design and the consensus; orange: stabilizing mutations introduced in the armadillo consensus design [10]; yellow: hydrophobic core positions.

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Designed repeat protein families.

Family	Family Repeat length	No. of rep ^a	Caps ^b	Architecture ^c	No. of res ^d
ank	33	3	des	a sol	168
TPR	34	4	ou	a sol	151
arm	42	4	des	a sol	252
HEAT	42	4	des	a sol	260
LRR	24	7	adapt	α/β sol	265
WD_6r	40	9	ou	β tor	259
WD_7r	40	7	ou	β tor	301
WD_8r	40	8	ou	β tor	346

Capping repeats, designed (des), adapted from native (adapt) or not present (no).

 c Solenoid (sol) or toroid (tor).

 $d_{
m Number}$ of residues.

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Ankyrin 6 TPR 6 Armadillo 8 HFAT 8	، م م	9	Monomer	Aggregates ^b	Folded ^c	Designs tested Expressed Soluble Monomer a Aggregates b Folded c Monomer T_{m}^{d} (°C) Yield e (mg/l) Crystal structures	Yield ^e (mg/l)	Crystal structures
rPR 6 Armadillo 8 HFAT 8	0		9		9	>95	80	4
Armadillo 8 HEAT 8	c	9	4	I	Sf	57, > 95	60	
HEAT 8	×	7	7		ю	>95	30	1
	8	5	1		ю	>95	30	
LRR 2	2	2	2	2	2	78	30	7
WD40_6rep 3	2	2	2	I			I	
WD40_7rep 3	3	ю						
WD40_8rep 7	4	4	3	3	1^{f}		10	
^a Determined by AGF-MALS.								
\boldsymbol{b} Soluble aggregates/oligomers were also observed in addition to the monomers.	were also observe	ed in additi	on to the mono	mers.				

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⁶Proteins were considered folded if they displayed a cooperative transition (sigmoidal shape) or no transition (signal loss of <20% at 95 °C) and if the CD spectrum at 25 °C corresponded to the expected secondary structure.

 d Midpoint of transition in thermal denaturation for the folded monomeric proteins within the family.

^eMilligrams of protein per liter of culture, as determined after ion metal affinity chromatography purification and dialysis. No optimization for protein expression was attempted.

 $f_{\rm Folded}$ protein forms a dimer. Only one TPR design is dimeric.