Computationally Designed Armadillo Repeat Proteins for Modular Peptide Recognition

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http://dx.doi.org/10.1016/j.jmb.2016.09.012
Edited by Amy Keating

Abstract

Armadillo repeat proteins (ArmRPs) recognize their target peptide in extended conformation and bind, in a first approximation, two residues per repeat. Thus, they may form the basis for building a modular system, in which each repeat is complementary to a piece of the target peptide. Accordingly, preselected repeats could be assembled into specific binding proteins on demand and thereby avoid the traditional generation of every new binding molecule by an independent selection from a library. Stacked armadillo repeats, each consisting of 42 aa arranged in three α-helices, build an elongated superhelical structure. Here, we analyzed the curvature variations in natural ArmRPs and identified a repeat pair from yeast importin-α as having the optimal curvature geometry that is complementary to a peptide over its whole length. We employed a symmetric in silico design to obtain a uniform sequence for a stackable repeat while maintaining the desired curvature geometry. Computationally designed ArmRPs (dArmRPs) had to be stabilized by mutations to remove regions of higher flexibility, which were identified by molecular dynamics simulations in explicit solvent. Using an N-capping repeat from the consensus-design approach, two different crystal structures of dArmRP were determined. Although the experimental structures of dArmRP deviated from the designed curvature, the insertion of the most conserved binding pockets of natural ArmRPs onto the surface of dArmRPs resulted in binders against the expected peptide with low nanomolar affinities, similar to the binders from the consensus-design series.

Introduction

Specific protein recognition is essential for many physiological processes and forms the basis of a number of procedures routinely used in research, diagnostics, and therapeutics. Still, the generation of binding reagents is time-consuming and has to be carried out for each desired target individually, either for monoclonal antibodies by immunization or for recombinant antibodies and alternative binding scaffolds by selection from a suitable library. A modular recognition of targets would allow the use of parts of the same binding surface in multiple contexts and speed up research and development by reducing design and selection steps. The complex features of globular protein surfaces usually prevent modular binding, but peptides are ideal targets for specific recognition by defined units, as in their extended form, they constitute regularly spaced structural features. As “peptides”, we refer not only to short stretches of amino acids but also to unstructured regions of proteins, such as termini, loops, or linkers between domains. Peptide–protein interactions are found in many highly dynamic cellular networks involved in signaling, regulation, and protein trafficking [1,2] and they represent about 15–40% of all
interactions in the cell [3]. For many applications involving recognition of such proteins or epitopes on denatured or digested forms of folded proteins, a general peptide-binding scaffold will be particularly valuable, if it can provide a modular and specific recognition of the peptide primary sequence.

Among the peptide-binding scaffolds, armadillo repeat proteins (ArmRPs) were found to form a stable framework with 4–12 repeats and provide a constant binding mode for peptides in an extended conformation, suitable for the generation of specific modular peptide binders (reviewed in Ref. [4]). Both natural (nArmRPs) and computationally designed ArmRPs (dArmRPs) consist of several internal repeats, which are stacked in tandem on each other to form an elongated α-solenoid protein with a continuous hydrophobic core and specialized capping repeats at the N- and C-termini. Each armadillo repeat unit, composed of 42 aa that fold into three α-helices (H1, H2, and H3) in a spiral staircase, binds approximately two consecutive amino acids of the peptide. This is achieved by a conserved asparagine residue, N 37 (superscripted numbers refer to the positions within the repeat) on each repeat, which fixes the peptide backbone by binding to every second peptide bond through bidentate hydrogen bonds, and by several surface residues forming pockets that interact with the peptide side chains [5].

In both major subfamilies of nArmRPs, importin-α and β-catenin, the peptides are bound in an antiparallel orientation (N- to C-terminal directions of protein and peptide run in opposite directions). The binding groove is built by a parallel arrangement of helix H3 of each repeat. In nArmRPs, the conserved binding mode is limited to three consecutive repeats (Fig. 1a) since the curvature is not constant across the entire protein. Thereby, the peptide units fall out of register with the armadillo repeats. In importin-α, two separated, negatively charged binding sites (major and minor binding site) are formed by repeats B-D and F-H, respectively, and can bind a bipartite nuclear localization sequence (NLS), with the typical sequence KR10-12K+x+ (“+” denoting any positively charged residue [6]). Therein, the two positively charged residue clusters are separated by a linker of 10–12 aa. In contrast to importin-α, β-catenin has only one binding site, which is itself positively charged, and the conserved binding is restricted to an area between repeats E-I.

This limited conserved binding of consecutive repeats in nArmRPs can be well explained by different curvatures found between neighboring repeat pairs. The analysis of nArmRP has shown that the sequence similarity between repeat units reaches only about 30%, and most differences—in length and residue composition—are found in the loops connecting the more conserved α-helices [7]. This low similarity is reflected in structural differences between repeats and, accordingly, in curvature variations between repeat pairs. In order to obtain a modular peptide-binding scaffold, curvature should be uniform over the whole protein and fit to the register given by the unit length of the peptide backbone.

Fig. 1. Peptide distance analysis of nArmRPs of the importin-α family. (a) Detailed view of the major binding site, composed of repeats B-E shown as gray cylinders for yeast importin-α (PDB ID: 1BK6) with bound NLS peptide (green), making six backbone hydrogen bonds (yellow dashed lines) with the conserved Asn residues (orange). Interaction residues of importin-α with peptide side chains are shown in yellow. Ionic interactions are indicated by blue dashed lines. The Ca(P/P+2) distance (red dashed lines) is measured between the Ca atoms (red spheres) of the peptide residues bound by Asn 37. (b) Summary of predicted Ca(P/P+2) distances found in repeat pair models of importin-α (BC to HI) distinguished by organism (calculated as described in Supplementary Fig. S1). N- and C-terminal repeats were excluded (repeat A and J). Upper and lower Ca(P/P+2) distances needed for continuous modular binding are indicated by black dashed lines. (PDB IDs: human: 2JDQ, 3FEX, 3FEY, and 3TJ3; mouse: 1EJL, 1EJY, 1IAL, 1IQ1, 1PJMN, 1Q1S, 1Q1T, 1Y2A, 2C1M, 3BTR, 3KND, 3L3Q, 3OQS, 3O5U, 3RZ9, 3RZX, 3TPM, 3UKW, 3UKX, 3UKZ, 3UL0, 3UL1, 3UVU, 3VE6, and 4HTV; yeast : 1BK6, 1EE4, 1EE5, 1UN0, and 2C1T).
A consensus-based design approach, which had been applied for other repeat proteins [8] and also for ArmRPs [5,9,10] (named consensus ArmRPs or cArmRPs), is expected to yield a uniform curvature, although it may not necessarily match the exact geometry desired for binding the dipeptide units. Therefore, an in silico design approach was developed in this work, based on a geometrically optimal curvature template. Using this template, the relative orientations between subsequent repeats were extracted and imposed as symmetric modeling constraints during backbone and side-chain sampling simulations using the Rosetta software suite [11]. Similar design protocols have been used for the computational design of repeat proteins, first with sequence and structural information obtained from natural repeat protein families [12,13] and then for de novo designed repeat proteins with open [14] and closed [15] architectures. Using such approaches, typically >50% of the designed constructs can be expressed as soluble, folded, and monomeric proteins, and determined structures agree well with the design models (typical RMSD of Ca atoms = 0.5–2.5 Å).

Next, protein regions of higher flexibility were assessed by molecular dynamics (MD) simulations, and more stable variants were engineered. Additional N-cap engineering allowed us to obtain an X-ray structure of a dArmRP, which resembled the original template model with an RMSD of about 1.9 Å. However, we found a deviation from the intended curvature. Nonetheless, surface modifications that had been grafted from nArmRPs–peptide complexes enabled the generation of binders against positively charged peptides with low nM affinities.

The structure of the dArmRP–(RR)_5 peptide complex revealed that the peptide is bound in an antiparallel orientation along the dArmRP binding surface as intended. This result demonstrates that although the exact desired curvature was not achieved, binding in a modular manner was still achieved for three consecutive repeats in the case of the dArmRP scaffold.

Results

Superhelical curvature of nArmRPs

For the development of a modular ArmRP, the superhelical curvature of the protein must match the distances found in the peptide in its bound conformation across many peptide units, that is, the location of each protein repeat must be exactly in register with the peptide bonds. To analyze this correlation between the peptide and repeat geometry, we described the superhelical fold of ArmRPs using helical symmetry parameters. We characterized the radius (r), rise (h), and angle (2·Ω), which together describe the positions of internal repeats (at its center of mass (CoM)) on a helix around a central axis (Supplementary Fig. S1). In the conserved binding mechanism, exemplified by the major binding site of yeast importin-α [16] (Fig. 1a), the position of every second peptide bond is defined through the interaction of a double hydrogen bond with N^32 in each armadillo repeat. Accordingly, a given ArmRP geometry provides a defined distance between a Ca atom of an amino acid (P) of the bound peptide and the Ca atom of two amino acids C-terminal to it (P+2).

When the peptide is bound in an extended conformation, the Ca(P/P+2) distance should be 6.7–7.0 Å (Fig. 1a). This Ca(P/P+2) distance is observed for bound peptides of importin-α, which assume an extended conformation within the major and minor binding site. This distance was also predicted by calculations and modeling of a peptide in relaxed β-strand conformation, including favored rotation angles and bond lengths. Note that the distance between two neighboring Ca atoms is constant at 3.8 Å because of the rigidity of the peptide bond, and because of the tetrahedral angle linking to units, the Ca(P/P+2) distance could maximally reach 7.0 Å.

In total, 36 peptide-bound structures of importin-α, which has 10 overall repeats, were analyzed geometrically by applying helical symmetry parameterization using Rosetta (Supplementary Fig. S1). Repeat pairs containing N- or C-terminal capping repeats were excluded from the analysis because of larger variations in sequence length and composition compared to internal repeats. It was found that only the repeat pair CD (from mouse and human importin-α) and GH (from yeast) display a curvature in relaxed β-strand conformation, including favored rotation angles and bond lengths. Note that this part acts like a linker, as described above for the bipartite NLS binding to importin-α.

Ideal ArmRP curvature for modular peptide binding

For the design of ArmRP possessing Ca(P/P+2) distances suitable for peptide binding, the GH repeat geometry was chosen over the CD repeats since GH repeats have lower Ca(P/P+2) distances and were considered to be more generic, as short distances were observed in all structures. Finally, we focused on the GH repeat geometry originating from yeast importin-α with optimal Ca(P/P+2) distances between 6.5 and 6.8 Å. To obtain a uniform multi-repeat curvature template with 12 repeats according to the GH repeat pair geometry (and to enforce a small Ca(P/P+2) distance), we superimposed repeat-wise
the copies of the GH repeat pair [Protein Data Bank (PDB) ID: 1EE4 [17]] (Supplementary Fig. S2). The GH backbone model provided an idealized ArmRP curvature compatible with binding a peptide over the whole length. The optimal curvature is characterized by a large radius ($r = 15.7$ Å), a small rise ($h = 6.2$ Å), and an intermediate angle ($2\cdot\Omega = 29.3^\circ$) in comparison to other curvatures found in yeast importin-α (Fig. 2a and Supplementary Fig. S2D). With these curvature parameters, a bound peptide is expected to have a Ca(P/P+2)-distance of 6.5 Å.

**In silico design**

In silico design was used to find an optimal amino acid sequence for the dArmRP that would support the desired curvature. Internal repeats were constrained to adopt the same primary sequence, side-chain, and backbone conformations by using symmetric sequence design and conformation sampling during all modeling moves. Symmetric structure prediction [18] and design have been used extensively in Rosetta [11], yielding atomic-accuracy.
predictions for large homomeric oligomers, designed cage-like assemblies, or repeat proteins [12–15, 19–21]. Our calculations were restricted to three internal repeats and two capping repeats. Since the modeled subunits are identical, the repeat protein could be extended indefinitely. We additionally enforced the N37 residues, because they are crucial for binding the peptide in each repeat. All other residues in the ArmRP were allowed to be mutated (to all amino acids except cysteine) using the Rosetta all-atom energy function (score12), which is dominated by Lennard-Jones, hydrogen-bonding, and implicit-solvent interactions [18].

Capping repeats were based on the computationally designed internal repeat. This was achieved by exchanging exposed hydrophobic residues in the capping repeats to hydrophilic ones (N-cap: A12E, P15Q, L19W, V27T, and A34Q; C-cap: V8E, V17E, L20Q, A25Q, A36E, and A39N), as had been done for designed ankyrin repeat proteins and dArmRPs from the consensus-design series [10].

Several positions in internal and capping repeats did not converge to a unique identity. Frequency in multiple sequence alignments and helical propensity were used to select residues for these positions. In the Rosetta models, positions 14, 15, 17, and 32 in the internal repeat (D-type); positions 17 and 32 for the N-cap; and positions 14, 15, and 37 for the C-cap allowed for several alternative residues (Fig. 3 and Supplementary Fig. S3). In the N-cap, Trp 19 was introduced to determine the protein concentration by UV absorbance at 280 nm. F38 was preserved in the C-cap as observed in nArmRPs, while the neighboring residues were redesigned using Rosetta.

The Rosetta models were analyzed by MD simulations, which revealed the instability of helix H3, probably due to the introduction of Ser residues with unfavorable helical propensity at positions 33 and 36. These positions were replaced by Ala residues, and an increase of stability was observed in MD simulations.

**Experimental validation of dArmRPs**

We systematically tested nine combinations of in silico designed variants experimentally (Supplementary Table S1). Each construct consisted of an N-terminal capping repeat, four internal repeats, and a C-terminal cap. Open reading frames were initially assembled as reported previously [5], but a faster one-day multi-fragment ligation assembly protocol could be developed (Supplementary Fig. S4). Proteins were purified by a single step of immobilized metal-ion affinity chromatography with yields of up to 80 mg per 1 L of Escherichia coli culture. Purified proteins were tested for their biophysical properties, that is, monomeric behavior, amount of secondary structure, accessibility of the hydrophobic core, and chemical stability.

Construct $N_V(D_{SPVA})_4C_{PAF}$ was identified as monomeric and folded, with lower 1-anilino-8-naphthalenesulphonate (ANS) binding, higher melting temperature, and, importantly, more cooperative unfolding compared to other variants, as measured by GdnHCl-induced unfolding. This construct was therefore chosen as the basis for further engineering (Fig. 4 and Supplementary Figs. S3, S5, and S6). This variant contains N- and C-terminal capping repeats of type NV and C_{PAF}, respectively, and four internal repeats of type D_{SPVA} (subscripts describe the variable residues that were experimentally found to be superior (see below), cf. Fig. 3 for sequence positions). For simplicity, $N_V(D_{SPVA})_4C_{PAF}$ will be named computationally designed ArmRP version 0 (CAR0). Additionally, CAR0...
was also identified by 1D $^1$H NMR experiments as a promising candidate, and 2D [$^{15}$N,$^1$H]-heteronuclear single quantum coherence spectra confirmed that CAR0 is stable and structured (Supplementary Fig. S7).

### Stabilization of dArmRP for structure determination

#### Stabilization by MD simulations

Several attempts to obtain crystals of construct N$_i$(DSPVA)$_x$CPAF with 1–10 internal repeats failed, although all proteins had favorable biophysical properties (Supplementary Fig. S8). Therefore, the Rosetta model of CAR0 was used as a starting structure for multiple explicit solvent MD runs (of 0.5 to 2 μs each) to identify regions with high flexibility and to stabilize them, as described before [9]. The MD trajectories showed that the overall fold of model CAR0 was preserved, with an average root mean square fluctuation (RMSF) of 0.4 Å for the C$_\alpha$ carbon atoms (the RMSD is given in Supplementary Table ST2). Higher conformational instability was observed for the caps with average RMSF values between 0.9 and 1.0 Å. The plasticity of the loop residues was higher than the helix residues with RMSF values of 0.96 Å and 0.74 Å, respectively.

On the basis of the RMSF values calculated along the MD sampling of CAR0, six mutations were introduced in each internal repeat to reduce fluctuations, resulting in the internal repeat Dq (Fig. 3). An additional set of independent simulations (of 0.5 to 2 μs each) were started from the mutated protein (with Dq internal repeats) and showed lower RMSF profiles than CAR0. Guided by the MD simulation results, the full-length construct CAR1 (N$_i$(Dq)$_x$CPAF) was produced and yielded a monomeric and well-folded structure.

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**Fig. 4.** Biophysical characterization of dArmRPs. (a) SEC (normalized absorption at 230 nm) and MALS (dots) of the proteins. Elution volumes of bovine serum albumin (MW: 66 kDa) and carbonic anhydrase (MW: 29 kDa) are indicated by dashed lines and were used as molecular weight standards. (b) ANS fluorescence spectra. Introduction of the Y III-cap resulted in ANS signals similar to the reference proteins, shown as horizontal dashed lines, indicating the highest ANS signal observed in the spectra of consensus-based proteins Y$_{IM}A_4$ and Y$_{IM}A_5$ [10]. (c) CD spectra of all proteins are shown and expressed as the MRE. (d) Normalized temperature-induced unfolding of designed proteins (dots) with fits (lines). (e) Normalized GdnHCl-induced unfolding of dArmRPs (dots) with fits (lines).
Table 1. Biophysical properties of dArmRPs

<table>
<thead>
<tr>
<th>Constructs</th>
<th>Short name</th>
<th>Residues</th>
<th>MWcalc (kDa)</th>
<th>MWobs (kDa)</th>
<th>OS</th>
<th>MWobs/cal</th>
<th>CD222 (MRE)</th>
<th>Tm (°C)</th>
<th>CDGdnHCl (M)</th>
<th>Kₐ (nM)</th>
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<tbody>
<tr>
<td>Nᵥ(Dᵥ(SPVA))ᵥCᵥPAF</td>
<td>CAR0</td>
<td>251</td>
<td>5.3</td>
<td>26.3</td>
<td>30.4</td>
<td>mono</td>
<td>1</td>
<td>−15,121</td>
<td>93 ± 1.8</td>
<td>2.2</td>
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<tr>
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<td>CAR1</td>
<td>251</td>
<td>5</td>
<td>26.3</td>
<td>31.9</td>
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<td>26.3</td>
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<tr>
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<td>25.2</td>
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<td>73 ± 0.3</td>
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n.d.: not determined.

* a dArmRP: N-cap (e.g., Nᵥ and Yᵥ), C-cap (e.g., CᵥPAF and Aᵥ), and internal repeats (e.g., Dᵥ(SPVA), Dᵥ, or Dᵥ(V₁)).

b The number of residues includes the MRGSH₆GS tag; all constructs consist of six repeats including capping repeats.

c Isoelectric point (pI) calculated from the sequence.

d Molecular weight calculated from the sequence.

e Observed molecular weight as determined by SEC.

f Oligomeric state (OS) measured by MALDI. Mono: monomeric state; Mix: equilibrium between monomer and dimer.

g Ratio between observed (by MALDI), and molecular weight calculated from the sequence (MWobs/calc).

h MRE at 222 nm expressed as deg·cm²/dmol.

i Transition midpoint (Tm) observed in thermal denaturation measured by CD (approximations).

j Midpoint of transition in GdnHCl-induced denaturation measured by CD.

k Equilibrium dissociation constant against (peptide)-sfGFP determined by fluorescence anisotropy.

l Consensus-based ArmRP. M refers to the consensus-based internal repeat M reported by Alfarano et al. [9].

m Data from Hansen et al. [31].
protein (Fig. 4). In comparison to CAR0, the stability increased by 0.6 M GdnHCl (Table 1). However, the increase in stability during chemical unfolding did not correlate with thermal unfolding data. We attribute this fact to an unusually high thermal (but not chemical) stability of the internal repeats D_{SPVA}. To guide the stability improvement of the dArmRP constructs, we relied on chemical denaturation data, since this procedure, in contrast to thermal denaturation, was able to fully unfold the proteins (as judged by CD spectroscopy) and showed a clear, cooperative transition. Crystallization attempts of CAR1 were, however, without success.

Replacement of the N-cap for crystallization

Crystal structures of dArmRPs were so far only determined for consensus-based ArmRPs with rationally designed N-caps (named Y_{III} and Y_{III}) [10]. In order to analyze the effect of the Y_{III} and Y_{III} capping repeat on dArmRPs in terms of biophysical properties and their ability to form crystals, protein CAR2 (Y_{III}(Dq)_{4}CPAF) was produced. The introduction of the Y_{III} cap increased the stability of CAR2 by 0.2 M GdnHCl, while thermal stability remained nearly identical (Fig. 4 and Table 1). Although the level of ANS binding increased compared to CAR1, it is similar to the well-folded, consensus-based ArmRP Y_{III}(M_{4}A_{II}) [10]. Overall, protein stability could be increased from CAR0 to CAR2 by a shift of 0.8 M GdnHCl in denaturation midpoint, which resembles the most stable dArmRP so far.

Structure of dArmRP CAR2

CAR2 crystallized at pH 5.5 in 0.1 M sodium acetate, 0.3 M sodium cacodylate, and 25% polyethylene glycol (PEG) 2K MME and diffracted to 2.0 Å resolution (Table 2). The structure was determined by molecular replacement. The two molecules in the asymmetric unit of CAR2 are aligned front to front by their C-terminal concave binding sites, burying a large surface area of 1682 Å^2 (Fig. 5). The right-handed superhelical structure of each molecule has an overall dimension of 60 x 30 x 20 Å. The lowest temperature factors were observed for the internal repeats (\(<B_{\text{nc-cap}}\) = 33.48 Å^2, \(<B_{\text{internal}}\) = 27.09 Å^2, and \(<B_{\text{C-cap}}\) = 42.30 Å^2), as previously observed for consensus-design ArmRPs [10,23] and other \(\alpha\)-solenoïd proteins [24–26] (Table 3). The largest temperature factors were found in the C-terminal capping repeat, especially within the loop connecting helices 2 and 3, which also displays no crystal contacts.

Structural comparison of dArmRPs

Both molecules from the asymmetric unit showed rather high structural variations (Fig. 5b). A comparison of chains A and B of CAR2 resulted in an RMSD of 1.1 Å [Ca residues from the N-cap (L13) to the C-cap (A247)]. The comparison of each individual internal repeat to the corresponding one in the chain B from the asymmetric unit revealed that they superimpose with an RMSD of 0.5 Å. The major exception is the loop connecting helices 2 and 3 in the internal repeat 3 of chain B. However, these different loop conformations do not explain the large RMSD for the superposition of the whole chains, which stems from small curvature variations between the repeat pairs.

Comparison of the experimental CAR2 structure with the designed model

The structure of CAR2 resembled the originally designed Rosetta-based model with RMSD of 1.8 Å ± 0.2 Å (Ca of four internal repeats, averaged over both molecules of the asymmetric unit). Although this value is usually acceptable for a general design approach, the analysis of the curvatures indicated a significant deviation from the initial design (Fig. 2). The model CAR0, which was obtained by Rosetta re-packing based on the curvature of the GH repeat from importin-\(\alpha\) (PDB ID: 1EE4), shows parameters (rise, radius, and angle) that result in a small Ca(P/P+2) distance of a modeled peptide. Accordingly, the overall shape can be described as a short and wide cylinder with a medium-sized angle between neighboring repeats (Fig. 2a). The structure of CAR2 deviates from the model curvature and can be described as a tall (large rise) and thin (small radius) cylinder with a large angle between neighboring repeats (Fig. 2b). Accordingly, we found that the expected Ca(P/P+2) distances of 8.2 ± 0.8 Å were significantly larger in CAR2 than in the initial design [Ca(P/P+2) distance of CAR0: 6.4 Å]. This difference between Ca distances of 1.8 Å indicates that although the design approach resulted in stable and typical \(\alpha\)-solenoïd folded proteins, the curvature of the apo-CAR2 structure is not meeting the requirements for a perfectly modular peptide-binding scaffold.

In contrast to nArmRPs, the curvature characteristics of repeat pairs from dArmRPs (Fig. 2b and c) are more uniform, which is expected because of their identical sequence in each internal repeat. However, due to the observed structural difference between the molecules within the asymmetric unit, the uncertainties in the parameters in dArmRPs, for example, the Ca(P/P+2) distances with standard deviations of 0.8 Å, are still rather large.

Surface redesign for peptide binding

The strongest affinities of nArmRPs were reported to be in the range of 20 nM for importin-\(\alpha\) binding to NLS peptides [27,28]. From an initial version of a consensus ArmRP library, a binder to neurotensin was selected with a \(K_d\) of 7 μM [29], but the binding...
mode was different from the intended canonical binding [30]. More recently, picomolar affinities for the interaction between the cArmRP YIII(Dq)4CPAF and the (KR)4 peptide have been measured [31]. In contrast, no significant peptide binding was detected for CAR2. This is not surprising, as surface residues potentially involved in binding were neither selected nor engineered for binding yet.

Therefore, a design of the dArmRP binding site was undertaken, inspired by the crystal structures of ArmRPs in complex with the target peptide. Similar approaches have been used to graft binding pockets onto the scaffold of tetratricopeptide repeat domains (TPR) [32–34]. The analysis of 36 structures of nArmRPs binding to NLS peptides revealed two highly conserved binding pockets at the major (P2 and P3) or the minor binding site (P1’ and P2’) [4], both binding the side chain of positively charged amino acids (Lys and Arg). Although the major and minor binding sites are similar, we focused on the minor binding site. In nArmRPs, the binding pocket P1’ is formed mainly by residues D1, T4, and A40, while P2’ is formed by E30, W33, and T40* (the asterisk indicates a position in the preceding repeat). Although position 40 is located distantly from the center of the binding pocket P2’, T40* in P2’ is highly conserved and forms one hydrogen bond to the backbone oxygen of the bound peptide. In order to

<table>
<thead>
<tr>
<th>PDB ID</th>
<th>YIII(Dq)4CPAF = CAR2</th>
<th>YIII(Dq.V1)4CPAF = CAR2.V1</th>
</tr>
</thead>
<tbody>
<tr>
<td>4D4E</td>
<td>with MRGSH6-tag no peptide</td>
<td>without MRGSH6-tag with (RR)5 peptide</td>
</tr>
<tr>
<td>Crystallization condition</td>
<td>0.3 M Na-acetate pH 5.5</td>
<td>0.1 M Tris–HCl pH 8.5</td>
</tr>
<tr>
<td></td>
<td>0.1 M Na-acetate</td>
<td>0.2 M Mg-chloride</td>
</tr>
<tr>
<td></td>
<td>25% PEG 2K MME</td>
<td>10% PEG 1000</td>
</tr>
<tr>
<td></td>
<td>10% PEG 8000</td>
<td></td>
</tr>
</tbody>
</table>

Data statistics

| Cell parameters | a: 51.81 Å,  | a: 88.57 Å, |
|                | b: 68.67 Å,  | b: 51.73 Å, |
|                | c: 126.84 Å, | c: 107.61 Å, |
|                | α: 90.0°,    | α: 90.0°, |
|                | β: 90.0°,    | β: 90.16°, |
|                | γ: 90.0°     | γ: 90.0° |

Space group | P212121  | P21 |
Resolution range | 39.35–2.0 Å (2.11–2.0) | 20.0–2.1 Å (2.15–2.1) |
Number of molecules/ asymmetric unit | 2         | 4 |
Number of reflections

| Observed | 139,251 (16,131) | 241,736 (18,790) |
| Unique  | 27,662 (3234)    | 106,755 (8086)   |
| I/σ(I)  | 9.3 (2.3)        | 5.01 (1.37)      |
| Completeness (%) | 88.2 (72.3) | 95.7 (87.4) |
| Rmerge (%) | 8.6 (58.8) | 16.7 (150.9) |
| Multiplicity | 5.0 (5.0) | 2.26 (2.32) |

Refinement statistics

| Rcryst (%) | 18.75 | 20.09 |
| Rfree (%)  | 26.34 | 25.45 |
| Number of protein atoms | 3490 | 7552 |
| Number of waters | 250 | 238 |
| Number of hetero atoms | 18 | 22 |
| Number of chains | 2 | 4 |
| RMSD from ideal geometry
  | Bond length (Å) | 0.016 | 0.015 |
  | Bond angles (°) | 1.875 | 1.795 |
| B-values (Å²) | Wilson B | 31.8 | 21.6 |
|                | Average B  | 33.5 | 43.5 |
| Ramachandran Plot (%)
  | Residues in preferred regions | 97.85 | 98.56 |
  | Residues allowed regions | 1.94 | 1.44 |
  | Residues in disallowed regions | 0.22 | 0.00 |

Values in parentheses are for the highest resolution shell.

1. $R_{\text{cryst}} = \Sigma |F_{\text{obs}} - F_{\text{calc}}| / \Sigma F_{\text{obs}}$, where $F_{\text{obs}}$ and $F_{\text{calc}}$ are the observed and the calculated structural factors, respectively.

2. $R_{\text{free}}$ was calculated using 5% of the reflections similar to $R_{\text{cryst}}$. 

Table 2. Data and refinement statistics
allow the occupation of both pockets over several repeats, we decided to place Ser at position 40, which can be seen as a compromise between Ala and Thr, to accommodate the requirements for P1′ and P2′. Surface mutations were introduced in the newly produced proteins CAR2.V1–CAR2.V4 (sequences are given in Fig. 3) to mimic pockets P2′ alone or P1′ in combination with P2′. Specific binding to positively charged peptides [e.g., (KR)4 and (KR)5] was qualitatively detected by ELISA for all surface variants, whereas only background binding was observed for protein CAR2. High background binding to the (KR)-peptides is expected due to cross-reactivity of the detection antibodies as illustrated by the no-protein control (Fig. 6).

In order to quantify protein affinity, we performed fluorescence anisotropy assays. Affinities were determined toward four different positively charged peptides ((RR)5, (RR)4, (KR)5, and (KR)4), fused to sfGFP as fluorescence marker. These peptides were chosen because during pocket design, no absolute preference for one of them could be deduced: within the analyzed 36 structures, pockets P1′:P2′ were found to be occupied by K/R with a frequency of 82%/6% : 97%/3%, while the frequencies for P2′:P3 were 9%/91% : 44%/44%, respectively. The His tag of all the surface-engineered dArmRPs was proteolytically removed to ensure an accessible binding groove (referred to as, for example, CAR2.V1_nohis). The affinity of the parental construct CAR2_nohis could only be approximately determined, since its low affinity to all peptides (>10 μM) would require very high protein concentrations that might interfere with assay conditions.

All the surface-engineered proteins gave rise to affinities in the nanomolar range, depending on the peptides tested (Table 1). The tightest interaction was observed for CAR.V2_nohis toward (RR)5 (Kd: 2.2 ± 0.1 nM), which corresponds to an increase in affinity of at least 4500-fold compared to CAR2_nohis. All surface-engineered binders show an increased affinity if the peptide is prolonged [(RR)4 versus (RR)5 or (KR)4 versus (KR)5], as previously determined for consensus ArmRP [31]. The effect is more pronounced for the (KR)n peptides, where one additional (KR) unit leads to an 11- to 16-fold affinity increase for different proteins, while one (RR)-unit increases the affinity 4- to 6-fold. The additional mutation K29Q, introduced in CAR.V1_nohis yielding CAR.V2_nohis, was designed to prevent the charge neutralization of E30 (Fig. 7c), which might therefore contribute to side-chain binding. This change resulted in an affinity increase by a factor of approximately 10 toward all peptides. The introduction of the second binding pocket P1′ (mutation N1D and I4T) only sometimes increased the affinity, for example, CAR2.V3_nohis compared to the parental CAR2.V1_nohis binding to (KR)n peptides. However, in other instances, the affinity did not change much [e.g., for CAR2.V3_nohis

| Table 3. Average B-factors of armadillo repeats (all values given in Å²) |
|-----------------|--------|--------|--------|
| chain           | N-cap | Internal| C-cap  |
| Y_{II}(Dq)_4C_{PAF} = CAR2     |   A    | 36.9   | 25.1   | 29.3   |
| B               | 30.0  | 29.1   | 55.3   |
| Y_{III}(Dq,V1)_4C_{PAF} = CAR2.V1 |   A    | 51.1   | 44.2   | 58.5   |
| B               | 44.9  | 31.1   | 63.5   |
| E               | 35.7  | 30.1   | 48.4   |
| F               | 45.6  | 44.4   | 75.8   |

B-factors were calculated from backbone atoms.
binding to (RR)$_n$ peptides] or was even decreased (e.g., for CAR2.V4_nohis compared to CAR2.V2_nohis when binding to all peptides). Therefore, contributions of individual mutations are not generally additive, and this point will require further investigations.

The constructs CAR2.V1-V4_nohis were monomeric, as measured by multi-angle light scattering (MALS), and eluted, with the exception of CAR2.V4_nohis, as a single symmetric peak, similar to CAR2 (Fig. 6). Although the modification of the surfaces did not alter the secondary structure, as characterized by the mean residue ellipticity (MRE) in CD spectra, it resulted in a decrease of stability by 0.3–1.4 M GdnHCl or by 10–22 °C (change of denaturation midpoints; Table 1).

**Structure of CAR2.V1 complexed with peptide (RR)$_5$**

For the structural investigation of the peptide–protein interactions, surface variants were set up for crystallization with peptides (KR)$_5$ or (RR)$_5$. A structure of CAR2.V1 in complex with an (RR)$_5$ peptide was determined at 2.1 Å resolution by molecular replacement using the previously determined crystal structure of CAR2.

During the refinement, a poly-arginine peptide was modeled into the density observed at the binding site, proving that each protein is binding one peptide. From the bound peptides, only 9 aa were resolved (Arg1–9 in chain A and C, and Arg2–10 in chain B and D). Additional density could be filled with single Arg residues, indicating that alternative binding conformations are possible. Two molecules of the CAR2.V1 structure are aligned front-to-front with a parallel orientation to each other (Fig. 8a). With this front-to-front orientation of two ArmRPs, the N- and C-termini of two antiparallel-bound peptides bound along the inner binding site are positioned closely to one another. Analogous to the CAR2 structure, the lowest temperature factors for CAR2.V1 were...
ArmRPs for Modular Peptide Recognition

Fig. 7. (legend on next page)
observed for the internal repeats, while the highest ones were seen for the C-cap (Table 3). The dArmRP molecules within the asymmetric unit are nearly identical (RMSD of 0.3 Å for all Ca atoms from residue 13 to 248), but the peptides differ significantly (Fig. 8b). Mutations introduced on the surface had no large impact on the overall structure. Thus, structures of CAR2 and CAR2.V1 superimpose with an RMSD of 1.1 Å (Ca of molecules of CAR2 superimposed on chain A of CAR2.+V1). This relatively large RMSD is a consequence of small structural changes in loops between helices 2 and 3 that are propagated along the solenoid. The structure of CAR2.V1 deviates slightly less than CAR2 from the curvature of the original model (RMSD of about 1.3 Å based on the Ca atoms of four internal repeats of CAR0). CAR2.V1 can be described analogously as a rather tall (large rise) and thin (small radius) cylinder with a large angle between neighboring repeats and with Ca(P/P + 2)-distances of 7.5 ± 0.2 Å (Fig. 2c).

Despite the enlarged Ca(P/P + 2) distance based on the protein curvature, peptides are bound along the designed binding surface in an antiparallel orientation and are fixed by several bidentate hydrogen bonds to asparagine residues (N37) on the surface (Fig. 7b), as intended in the original design and observed in nArmRPs. Superposition of all molecules revealed a highly conserved backbone of the bound peptide. Conformational fluctuations of the peptide backbone and side chains were observed toward the C termini of the peptide; thus, the highly conserved binding of the side chain is observed only for Arg2, Arg4, and Arg6 in pocket P1' and for Arg7 and Arg9 in pocket P2', respectively. The increased conformational space sampled toward the ends of the peptide is consistent with its increased temperature factor and with a reduced number (e.g., chain A and E) or non-ideal geometry (chain B and F) of bidentate hydrogen bonds formed by N37 to the backbone of the bound peptide (Fig. 7b). The measurement of the Ca distances of the bound peptide revealed Ca(P/P + 2) distances of 6.6 ± 0.2 Å (measured for the distances among Arg3, Arg5, Arg7, and Arg9 of the peptide). Thus, binding of the poly-arginine peptide to protein CAR2.V1 with its non-optimal curvature was only possible by reducing the number of repeats involved in modular binding.

Despite the deviation from the optimal binding mechanism, single engineered pockets do bind the peptide side chains. As expected from the design, binding pockets are located between neighboring internal repeats and are only partially formed if capping repeats are involved. Accordingly, the introduced binding pockets P2' make interactions with the positively charged side chains of the peptide (Arg5, Arg7, and Arg9). The binding mechanism of pocket P2' is highly conserved (Fig. 7c). The mutation N30E allows the formation of a salt bridge between E30-OE1 or OE2 and the peptide side chain Arg-NH1 or NH2 (Arg5, Arg7, and Arg9; Arg3 is excluded because no negatively charged amino acid is located at position 30 in the C-cap). Although the conformation of the introduced W33* is more variable, the mutation A35W allows the formation of a cation–π interaction [35]. This is achieved with W33* either in an upright (e.g., Trp117) or in a flattened conformation (e.g., Trp159). Although no binding pockets were designed at position P1' (required to bind Arg2, 4, 6, and 8) in CAR2.V1, residues G41*, N1, and S40 allow the formation of a four-hydrogen-bond network and fix the peptide to the surface. G41*-O interacts with R3-NH1, N1-O interacts with R3-NH2, and S40 makes two hydrogen bonds, namely S40-O with R3-NH2, and S40-OG with R3-NE (Fig. 7d). Thus, the mutation of A40*S seems to be beneficial for binding arginines in pockets P1'. The mutations N1D and L4T additionally used for grafting P1' pockets in construct V3 and V4 did not always increase the affinity for (KR)n, or (RR)n, compared to the constructs missing these two mutations (V1 and V2; see Table 1). This indicates that the designed, complete binding pocket P1' in CAR2.V3 and V4 and the observed hydrogen network in CAR2.V1 and V2 that is made by wild-type residues and A40*S are expected to fix the peptide side chains with similar efficacy.

The comparison of apo- and holo-structures of dArmRPs revealed that binding to the poly-arginine

Fig. 7. Peptide-binding mode in CAR2.V1. (a) Superposition of poly-arginine peptides [(RR)n shown as blue sticks] on chain A shown in surface representation and colored according to electrostatic potential. Binding pockets P1' are occupied by residues Arg2, Arg4, Arg6, and partially Arg8, whereas rationally engineered binding pockets P2' are filled with Arg7, Arg9, and partially Arg5. N- and C-terminal residues of peptides are more flexible and not shown. (b) Modular binding mechanism of peptide backbone (salmon stick) of ArmRP chain B. Part of helices H3 are shown as cylinders. Bidentate hydrogen bonds to Asn37*, and measured Ca(P/P + 2) distances are indicated as yellow and red dashed lines, respectively. Conserved modular backbone binding was observed for residues Arg5 and Arg7. Arg3 and Arg9 are bound in a less conserved manner, indicated by the increased length or complete absence of hydrogen bonds in internal repeat 1 or 4 of chain A–E. (c) Conserved binding mode in pocket P2'. Arginine residues are fixed by ionic interactions with Glu30* (blue dashed lines) and cation–π interactions with Trp30. Lys29* (green) was mutated to Gin in CAR2.V2 and CAR2.V4 to remove the charge neutralization of adjacent Gin30*. Plain numbers and superscripts refer to the numbering scheme in the PDB file and individual repeats, respectively. (d) Conserved binding mechanism in pocket P1' composed of four hydrogen bonds mediated by Gly41*, Asn1*, and Ser40* (* and # indicate positions in previous and following repeats, respectively).
peptide had only moderate effects on the overall protein curvature. While structural variations observed in CAR2 are still present in CAR2.V1, indicating that the overall curvature is similar, and a certain amount of flexibility is maintained in the scaffold, and the variations in the complex structure are slightly smaller. Nonetheless, the overall curvature of CAR2.V1 did not adapt upon binding to the peptide, and thus, the peptide did not induce a complete modular binding all along its length.

The holo-structure also explained the beneficial effect of the charge neutralization mutation K29Q in CAR2.V2 on affinity. K29 in CAR2.V1 is located on the binding surface close to binding pocket P2’ (see Fig. 7c). Thus, the positive charge of K29 would reduce the charge–charge interaction between the ligand arginine residue and E30. This effect is strengthened by the multiple appearance of this pocket in the repetitive binding molecule.

**Discussion**

To design an ArmRP with curvature geometry suitable for modular peptide binding, we applied a computational approach based on the multi-repeat model from yeast importin-α repeat pair GH. The introduction of six MD-based mutations within the
internal repeat and the replacement of the N-cap by the consensus-based Yres-cap were necessary to improve thermodynamic stability and obtain crystal structures of dArmRPs. The structural analysis revealed that dArmRPs deviate from the planned curvature that would be optimal for binding peptides. The Ca(P/P +2) distance is about 8.2 Å, which is significantly more than the desired binding distance of 6.7–7.0 Å. Nonetheless, modification of the binding surface of dArmRP resulted in binders with the ability to bind (RR)n and (RR)n peptides. Affinities as high as 2.2 nM were determined by fluorescence anisotropy (Table 1). Importantly, the crystal structure of the complex with (RR)n clearly shows that the side chains occupy the engineered cavities and thereby exclude non-specific electrostatic interactions as the main mode of binding, also consistent with the 1:1 stoichiometry deduced from the fluorescence anisotropy binding curve (Fig. 6e).

**Computationally designed armadillo scaffold**

The C_paf-cap of CAR0 was the only part of the *in silico* design, of which a structure could be obtained, without further modifications, and therefore it allows the estimation of precision of the design. A comparison of the C_paf-cap of the Rosetta model and its X-ray structures (CAR2 and CAR2.V1) superimposes with a low RMSD of 1.1 Å (based on the Ca atoms) and highlights the accuracy of the design (Fig. 9a). The largest difference between the model and the structures was observed in the loop connecting helices 2 and 3. This loop is rearranged and allowed H22 to bind into the groove between the last internal repeat and the C-cap. Thereby, H22 forms a hydrogen bond with the backbone of A24 and is involved in the interaction network of the hydrophobic core by forming contacts with residues L19, N24, and V27 of the C-cap and residues E25 and I28 of the internal repeat (Fig. 9b). Notably, the same loop conformation is observed between internal repeats, although H22 is replaced by D22 there. In addition, the C_paf-capping repeat has a nearly identical structure to the consensus-based Ares-capping repeat [10] (PDB ID: 4DB6) (Fig. 9c). Although they share only 63% sequence identity and 68% sequence similarity, the RMSD is 0.8 Å (based on the Ca residues of both C-caps).

**Stabilizing effect of the MD-based Dq internal repeat**

The structural stability of the CAR0 model was analyzed by MD simulations, and several mutations were introduced in the internal repeats (Dq) at positions of high RMSF. The mutations M3Q, L4I, V8I, E21D, K28I, and A32V (in CAR1) indeed increased the overall stability of this protein as confirmed experimentally (Table 1).

As armadillo repeats are largely defined by the conservation of hydrophobic core residues, and as the stability of the protein largely depends on the continuous hydrophobic core [36], the mutations in the core ([L4I, V8I, K28I, and A32V]) and intermediate region (M3Q) are expected to be essential for the overall increase in stability (Fig. 10). Among them, the side chain of Q3 is partially involved in the hydrophobic core, but its hydrophilic end is involved in three conserved hydrogen bonds (Fig. 10b). Methionine at position 3 cannot form these hydrogen bonds, and additional clashes are found for all possible rotamers sampled by Rosetta (data not shown).

The largest change in side-chain size was the mutation A32V. Here, several alternative residues (Ala, Leu, and Cys) were suggested by Rosetta. Leu showed no measurable advantage over Ala (Supplementary Fig. S5 and Supplementary Table ST1), and Cys was not further considered in order to avoid issues with disulfide bond formation. MD simulations with A32 revealed a small cavity, which was filled with Val, and this packing was confirmed in the CAR2 structure. In contrast, Leu, when modeled into this position, would clash with several other side chains of the core (Fig. 10a).

Analogously, Ile fits best at position 8, since it is involved in more van der Waals interactions compared to Val, and Leu would result in clashes (Fig. 10c). The unfavorable effects of L8 were tested experimentally, and indeed, protein stability was decreased by about 0.5 M GdnHCl (midpoints of denaturation; data not shown).

In position 4, Leu was replaced by Ile. In the structures of CAR2, Ile fills the hydrophobic core cavity without clashes, in contrast to Leu (Fig. 10d). Therefore, Ile residues at position 4 and 8 seem to be essential for the overall compactness and stability of the structure. At position 28, both residues (Lys and Ile) seem to be valid solutions to strengthen the protein core; however, Ile better fills the hydrophobic core without leaving any cavity.

**Curvature deviation from optimal template**

Structural analysis of CAR2 showed that the desired curvature was not obtained, and the corresponding Ca(P/P +2) distances are too long to match the bound peptide over a longer distance. For an efficient comparison of ArmRPs with different curvatures, we characterized the overall superhelical curvature by four parameters [rise, radius, angle, and Ca(P/P +2) distance]. The accuracy of the parameterization depends on how well the structure can be described with the applied symmetry, and several tests verified the validity of the parameterization: first, each parameterized structure from CAR2, consisting only of the backbone Ca atoms of two neighboring repeat pairs, superimposes with real structure pairs with a low RMSD of 0.5 Å ± 0.1 Å. Second, the Ca(P/P +2) distance measured on the parameterized structures does not deviate more than 0.1 Å from distances measured directly on the experimentally determined structures.
Fig. 9. Structural details of the computationally designed cap $C_{PAF}$. (a) Superposition of the $C_{PAF}$-cap from the Rosetta model (CAR0, gray) and the crystal structure (CAR2, magenta). Major differences are found in the loop region (light magenta). (b) Detailed view of the loop conformation mediated by residue His$^{22}$ (yellow). Conserved hydrogen bonds to residue Asn$^{24}$ (yellow dashed line) and several hydrophobic interactions (blue dashed lines) are found to be identical in the internal repeats Dq (shown in gray) involving residue Asp$^{22*}$ (orange). (c) Structural similarity of $C_{PAF}$ superimposed on the C-cap of consensus-based cap $A_{II}$ (green; PDB ID: 4DB6 [10]).

Fig. 10. Structural details of stabilizing mutations introduced by MD simulations in internal repeats Dq. (a) Mutations introduced in the internal repeats and the C-cap are shown in stick representation in the structure of CAR2. N- and C-cap are colored in orange and green, respectively. (b–d) Introduced residues in the Dq sequence are colored in green and are compared to either the modeled wild-type residue (D$_{spVA}$) or another mutation (yellow). Residues in close proximity are shown in gray stick representation (* and # indicate the positions in the preceding and following repeats, respectively). Hydrogen bonds are indicated by yellow dashed lines. Clashes observed in the model wild-type structures are shown with red cylinders. (b) Gln at position 3 makes three hydrogen bonds in contrast to Met. (c) Hydrophobic core mutation V8I fills the hydrophobic core completely (Ile shown in green). Mutation I8L (§: a mutation introduced in a CAR2 variant; data not shown) results in several clashes and a reduced protein stability. (d) Ile fits into the hydrophobic core at position 4 without clashes.
The impact of the Y III-capping repeat was additionally protein CAR0 (N V(DSPVA)4CPAF). In contrast to the tested on the original, computationally designed (Supplementary Fig. S9). Accordingly, YIII and Dq must have been expected in the Rosetta-designed model of CAR0. In contrast to CAR2, no crystals could be obtained for YIII(DSPVA)4CPAF, and thus, we cannot experimentally investigate the differences in interaction.

These results do not allow us to decide whether the capping repeat has an influence on the overall curvature or not. Nevertheless, several designed caps in consensus-based ArmRPs (YIII, YII, and AIII) [10] have shown that the overall curvature is not influenced by the capping repeats. So far, the only influence was observed for internal repeat pairs adjacent to a domain-swapped cap, as found in the early structures of consensus-based ArmRPs [10,23].

The impact of stabilizing mutations on the overall curvature of the protein is difficult to predict, but computational analysis can provide insights. We observed in the MD simulations that the mutations introduced in CAR1 induced a change of the overall curvature toward more unfavorable parameters for modular peptide binding, resulting in the curvature also observed in the experimentally determined structures of CAR2. Accordingly, the Ca(P/P+2) distance changed in the model of CAR1 from 6.4 Å to 7.6 Å during the simulations. We hypothesize that the larger side chains introduced when converting the DSPVA to the Dq repeat (L4I, V8I, and A32V) forced the protein to stretch from short and wide toward a longer but thinner shape. Among the introduced mutations, V32 seems to be a candidate to induce a curvature change, similar to what has been expected in the Rosetta-designed model of DPAAL with L32 (Supplementary Fig. S3). However, at this stage, these explanations remain largely speculative because structural information from the original CAR0 sequence is not available.

**Stabilization effect of the YIII-capping repeat with the Dq internal repeat**

Terminal capping repeats have been shown to be essential for the stability of repeat proteins, demonstrated, for example, in DARPins [22] and consensus-based ArmRPs [9,10]. For dArmRPs, the impact of the YIII-capping repeat was additionally tested on the original, computationally designed protein CAR0 (NIV(DSPVA)ATCPAF). In contrast to the stabilizing effect of YIII observed in CAR2 (Table 1), the addition of YIII slightly reduced the stability against the chemical denaturation of YIII(DSPVA)ATCPAF (Supplementary Fig. S9). Accordingly, YIII and Dq must have a favorable interface complementarity, which enhances the overall protein stability by 0.4 M GdnHCl in comparison to YII and DSPVA. In contrast to CAR2, no crystals could be obtained for YIII(DSPVA)ATCPAF, and thus, we cannot experimentally investigate the differences in interaction.

### Peptide binding by dArmRPs

The surface of CAR2 was modified and guided by structures of nArmRPs in complex with target peptides and by the knowledge from consensus ArmRP–(KR)n interactions [31]. The introduction of the binding pocket P2′ of importin-α into each internal repeat resulted in strong and specific binding molecules of positively charged peptides. The additional peptide-binding pocket P1′ (mutations N1D and I4T) increased the affinity only in some instances. One explanation for their small effect is the presence of a hydrogen bond network—present already in CAR2—which allows the fixation of the corresponding peptide side chains at similar positions as predicted for P1′ pockets, which therefore cannot contribute more binding energy. Nonetheless, the placement of a negatively charged residue as an anchor point in the P1′ pocket could still be a valid strategy to improve the binding of a positively charged side chain. Accordingly, D1 could be replaced by E1 to gain more flexibility or be shifted to position 41, as found in the pocket P2 of the major binding site of importin-α.

Our finding that the crystal structures of CAR2 and CAR2.V1 deviated from the expected modular binding scaffold is also reflected in the observed binding mode of the peptide. In the complex structure, dArmRP can bind a peptide, but only over a short stretch of approximately three repeats. Consequently, more flexibility is observed in the bound peptides toward their termini, indicating the loss of binding at the peptide ends.

In future design cycles, the structure should be further optimized in order to result in a curvature that will allow modular binding over more repeats. With the availability of an increasing number of crystal structures of dArmRPs, with or without bound peptide, more details about curvature and its impact
on peptide binding will emerge. For example, we observed that CAR2 has larger structural fluctuations, as deduced from the larger SDs in their curvature parameters. Binding of the peptide in CAR2.V1 seems to restrict the curvature to a state slightly closer to the desired one [reduced Ca(P/P+2) distance].

In the future, it needs to be investigated whether directed evolution experiments are capable of identifying mutations that alter the curvature to become compatible with modular binding over longer distance. Furthermore, knowledge from additional experimental structures of dArmRP, in combination with protocols for enhanced sampling by MD [37], will likely allow a manipulation of curvature accurate enough to obtain modular binding over wide distances.

Materials and Methods

General molecular biology methods

Unless stated otherwise, experiments were performed as described previously [5].

Gene assembly and protein expression

Full-length gene assembly for proteins containing an N-terminal capping repeat, several internal repeats, and a C-terminal capping repeat was performed as described previously [5] by using a step-by-step ligation of the modules digested by Bsal or BpiI or by using a single multi-fragment ligation step (described below), similar to what has been described elsewhere [38,39]. Exchange of capping repeats on full-length constructs was performed by PCR amplification using two overhang fragments (20 bp) encoding the new capping repeat and the template fragment, which is lacking the corresponding capping repeat. Full-length genes for binder CAR.V1–CAR.V4 were synthesized by GeneArt® (Life Technologies) and cloned with BamHI and HindIII into the pQE-derived vector.

Expression and purification for biophysical characterization and for crystallization was done as described previously [23]. Protein concentrations were determined by absorbance at 235 and 280 nm using molecular masses and extinction coefficients calculated with the tools available at the ExPASy proteomics server. Protein size and purity were assessed by 15% SDS-PAGE stained with Coomassie PhastGel Blue R-350 (GE Healthcare, Switzerland) and confirmed by mass spectroscopy.

Multi-fragment ligation assembly

Multi-fragment ligation (Supplementary Fig. S4) was used to assemble dArmRPs with 1–10 identical internal repeats in one step. For this purpose, single modules were amplified by PCR with primer pairs pQE30_for + pQE30_short_KpnI_r, pQE30_short_BamHI_f+pQE30_short_KpnI_r, or pQE30_short_BamHI_f+pQE30_rev from vectors containing the N-capping, internal, or C-capping repeats, respectively (primers are given in Supplementary Table S3). The extended overhang sequence upstream and downstream of the N- and C-capping repeat provides an important quality control to obtain full-length constructs after PCR amplification of the gel-purified multi-fragment ligation product. The corresponding PCR fragments were digested by Bsal, Bsal + BpiI, and BpiI and were purified by NucleoSpin columns (Macherey-Nagel). Ligation was performed in three steps: first, 0.5 μg of single-digested N-cap fragments were ligated with a fivefold excess of double-digested internal repeat fragments using 2.5 U of T4 DNA Ligase and were incubated for 15’ at room temperature. In the second step, 0.5 μg of digested C-cap fragment and 1 U of fresh ligase were added to the mixture after buffer adjustment according to the volume increase and were incubated for 30’. In the last step (optional), 0.5 μg of N- and C-cap fragments were added again to the mixture to increase the amount of full-length constructs. The buffer was adjusted according to volume increase and incubated for another 15’. To obtain constructs with the right number of internal repeats, the ligation mixture was heat-inactivated at 65 °C for 10’ prior to loading and was separated on a 1.5% agarose gel. The desired DNA bands were extracted and amplified by PCR with outer primers pQE30_for + pQE30_rev using 50 ng template DNA. Full-length fragments were inserted into cloning and expression vectors pQE30 or pPANK by BamHI and HindIII sites. Proper assembly of constructs was validated by colony PCR and DNA sequencing.

Model backbone template generation

A backbone model based only on the GH curvature from importin-α (PDB ID: 1EE4 [17]) was generated by iterative superposition of the repeat G from the GH repeat pair on the H repeat of a copy of the GH repeat (Supplementary Fig. S2). The GH–backbone template was then used to repack the protein by the Rosetta Program.

In silico protein design (Rosetta software)

Symmetric constraints were applied throughout the design trajectories. A single “master” ArmRP domain was designated arbitrarily, and all side-chain packing, minimization, and backbone minimization moves were done simultaneously on this master domain and all other domains in the system. Each move consisted of combinatorial side-chain design and conformational search, and backbone and side-chain minimization. An extended peptide, corresponding to the target peptide for binding, was maintained throughout the simulations in the preferred orientation observed in the yeast importin-α crystal structure (PDB ID: 1EE4 [17]). The Asn residues at position 37 that contact the peptide backbone were maintained in their native orientations, and the rigid-body orientation of the peptide–ArmRP complex was minimized during the simulation. Resulting models were analyzed for their peptide–backbone binding ability and structural integrity. Further symmetric substitutions were introduced on the surface of the ArmRP by adding salt bridges and polar groups to increase solubility.
Analysis of superhelical parameters

Superhelical parameters were determined by analyzing the geometry of internal repeat pairs using the generalized helix description, as it has been implemented in the make_symmdef_file.pl script from the Rosetta symmetry framework [18]. For input structures, we used the Cα atom coordinates from 41 residues of two consecutive internal repeats (the flexible residues at position 23 were excluded). Curvature parameters as depicted in Fig. 2 were first generated for each pair of internal repeats (M1:M2, M2:M3, M3:M4, and M4:M5) and for each molecule found within the asymmetric unit and were then averaged (Supplementary Fig. S1C). The angle 2·Ω (°) describes the angle between the centers of the mass of two consecutive internal repeats (Supplementary Fig. S1). The angle 2·Ω (°) was used to probe the packing of the designed hydrophobic core.

CD spectroscopy and unfolding curves

CD measurements were performed on a Jasco J-810 spectropolarimeter (Jasco, Japan) using a 10 μM protein solution in 50 mM Tris (pH 7.6) and 150 mM NaCl in a 0.5-mm cylindrical thermocuvette. CD spectra were recorded from 190 to 250 nm with a data pitch of 0.5 nm, a scan speed of 100 nm/min, a response time of 4 s, and a bandwidth of 1 nm. Each spectrum was recorded four times and was averaged. Measurements were performed at 20 °C. The CD signal was blank-corrected and converted to MRE. GdnHCl-induced denaturation measurements were performed after overnight incubation at 20 °C with increasing concentrations of GdnHCl (99.5% purity, Fluka), and the data were collected at 222 nm (data pitch = 1 s; response time = 4 s; bandwidth = 2 nm; and measured time = 45 s) and processed as described above. Heat denaturation curves were obtained by measuring the CD signal at 222 nm with temperature increasing from 20 to 92 °C using an external water bath (Julabo FS18; data pitch = 0.2 °C; heating rate = 1 °C/min; response time = 4 s; and bandwidth = 1 nm).

GdnHCl-induced denaturation data showed slopes of the pre-transition and post-transition phases that are either close to zero or not well defined, and thus, they were set to zero. Data were thus fitted to a two-state unfolding model without sloping baselines (Eq. (1)). Fits were only used to estimate the unfolding midpoint and not the other parameters.

\[
Y = \frac{y_f + y_u \times e^{-\Delta G_{m}/RT}}{1 + e^{-\Delta G_{m}/RT}}
\]

with \(\Delta G = \Delta G_{m} - m_{[\text{GdnHCl}]}\), where \(Y\) is the fraction of unfolded protein (expressed as normalized MRE); \(y_f\) and \(y_u\) are the signals for fully folded and fully denatured proteins, respectively; \(R\) is the universal gas constant, \(T\) is the temperature (298 K), \([\text{GdnHCl}]\) is the concentration of GdnHCl, and \([\text{GdnHCl}]_{1/2}\) is the GdnHCl concentration at the unfolding midpoint, which is obtained from \([\text{GdnHCl}]_{1/2}\) = \(\Delta G_{m}/RT\). For the denaturation plots, values of MRE (blank corrected) were normalized by setting the pre-transition values (folded) as 0 and the putative, completely unfolded protein (defined as MRE = 0) as 1.

Thermal denaturation curves were fitted to a two-state unfolding model with sloping baselines according to Jackson and Fersht [40] with Eqs. (2) and (3). Since the full reversibility and two-state nature of this system are questionable, all fits were only used to estimate the midpoint of thermal denaturation and not the other parameters:

\[
\Delta G = \frac{T_{m} - T}{T_{m}} \times \Delta H \times (T_{m} - T) \times \Delta C_{p} + T \times \Delta C_{p} \times \ln \frac{T_{m}}{T}
\]

\[
\text{MRE} = \frac{1}{1 + e^{-(y_{f} + m_{T} - y_{u} - m_{u} \times T) + y_{u} + m_{u} \times T}}
\]

Here, \(y_{f}\) and \(y_{u}\) are the y-axis intercepts of the lower and upper baseline, respectively; \(m_{T}\) and \(m_{u}\) are the slopes of the lower and upper baseline, respectively; \(T\) is the temperature, and \(T_{m}\) the midpoint of thermal denaturation; \(\Delta G\) and \(\Delta H\) are the free energy and enthalpy of unfolding, respectively, and \(\Delta C_{p}\) is the change in heat capacity at constant pressure.
NMR spectroscopy

All 1D $^1$H NMR spectra were recorded at 37 °C using 0.5 mM protein solutions in 50 mM phosphate buffer (pH 7.6) and 30 mM NaCl. 1D proton NMR spectra were recorded with water suppression by standard presaturation. To obtain proton–nitrogen correlation maps, standard $^{15}$N,$^1$H-heteronuclear single quantum coherence spectra were recorded on Bruker Avance 700 MHz NMR spectrometer with uniformly labeled $^{15}$N-labeled protein in 50 mM phosphate buffer (pH 7.6) and 150 mM NaCl [41]. Data were processed and inspected in Topspin 2.1.

Crystallization and structure determination

Sparse-matrix screens from Hampton Research (California) and Molecular Dimensions (Suffolk, UK) were used to identify the preliminary crystallization conditions in 96-well Cominig plates (Cominig Incorporated, New York) at 4 °C. A Phoenix crystallization robot (Art Robbins Instruments) was used to perform sitting-drop vapor-diffusion experiments. The protein solutions were filtered through a 0.22 μm Millex® filter (Millipore). Prior to crystallization, protein CAR2.V1 was supplemented with a 1.5-fold molar excess of (RR) peptide (the peptide was dissolved in water, and changed the volume of the sample by 1%). Proteins CAR2 and CAR2.V1_nohis were mixed with reservoir solutions at 1:1, 1:2, or 2:1 ratios (200–300 nl final volume) and at 1:1, 1.2, or 1.5 (300 nl), respectively, and the mixtures were equilibrated against 50 μl of reservoir solution at 4 °C. Table 2 summarizes all the crystallization conditions, data collection, and refinement statistics. Crystals for CAR2 and CAR2.V1 were washed three times in reservoir solution supplemented with 10% glycerol and 20% ethylene glycol, respectively, before being flash-cooled in liquid nitrogen. Using a Pilatus detector system on beam line X06DA at the Swiss Light Source (Paul Scherrer Institute, Villigen, Switzerland), data were collected and processed using the program XDS [42]. The CAR2 structure was solved by molecular replacement (software PHASER [43]) using a structure of another CAR2 variant, which had been determined by a poly-alanine search model that was created from the crystal structure of a consensus ArmRP (chain A from PDB ID: 4VSR [44]). CAR2.V1_nohis was determined by using CAR2 as search model. Refinement was done using programs PHENIX-Refine [45], REFMACS [46], and COOT [47]. Water molecules were added to well-defined difference electron density peaks at hydrogen bond distance from the protein. (RR) peptides were identified in the final electron density maps of CAR2.V1. The program PROCHECK [48] was used to validate the final structures, and PyMOL was used to generate figures [49].

Diffraction data of CAR2.V1 suggested an orthorhombic space group, and structure determination was possible in space group P2$_1$2$_1$2$_1$ with two molecules related by a 2-fold axis. In this setting, the non-crystallographic rotation axis was almost parallel to the c-axis. However, the refinement did not converge, and the $R_{max}$ value never dropped below 29.3%. Therefore, the symmetry restraints were relaxed and the structure was refined in space group P2$_1$ with four molecules in the asymmetric unit. All four chains show small but significant deviations in ligand binding, which confirms that the assignment of a monoclinic space group is correct, although the crystal lattice is almost orthorhombic. Further analysis of the diffraction data using an L-test [50] implemented in the program TRUNCATE [51] suggested pseudo-merohedral twinning with a twinning fraction of 49.5%. Only after taking twinning into account, the refinement converged at final $R_{cryst}$ and $R_{free}$ values of 20.19% and 25.53%, respectively.

MD simulations

Explicit solvent MD simulations were performed at constant temperature (310 K) and constant pressure (1 atm) using the v-rcalcus thermostat [52] and Berendsen pressure coupling [53]. The long-range electrostatic interactions were treated by the particle mesh Ewald method [54]. The van-der-Waals interactions were truncated at a cutoff of 9 Å. The LINCS algorithm [55] was used to fix the length of all bonds. Virtual sites were used for removing the fastest degrees of freedom, which allowed an integration time step of 5 fs. Structures were saved every 5 ps for analysis. The MD simulations were carried out using the Gromacs program [56] with the OPLS force field [57,58] and the TIP3P potential [59] for water molecules. The first 200 ns of each MD run were considered as equilibration time and were excluded from the RMSF calculation. The block average time in RMSF calculation was 2 ns. All the constructs were simulated in the absence of the His tag.

ELISA

A MaxiSorp plate (Nunc) was coated with NeutrAvidin (100 μl, 66 nM in phosphate buffered saline (PBS) overnight at 4 °C) and then blocked with PBS-TB (300 μl, 0.1% Tween, and 0.2% bovine serum albumin (BSA) in PBS) for 1 h at room temperature. The target peptides (expressed as pD-fusion [5,29] or chemically synthesized (JPT) (Supplementary Table ST4)) were immobilized via their biotin residues on NeutrAvidin (100 μl, 200 nM in PBS-TB). Buffers for binding and washing in all ELISA experiments were PBS-TB and PBS-T (300 μl, 0.1% Tween in PBS), respectively. Purified proteins (100 μl, 200 nM in PBS-TB) were incubated with the target for 1 h at 4 °C. Wells were washed three times with 300 μl of PBS-T before the detection of the proteins with a primary anti-RGS18 antibody (100 μl, 1:5000 dilution in PBS-TB for 45 min at 4 °C; Qiagen, Germany) and a secondary goat anti-mouse IgG alkaline phosphatase-conjugated antibody (100 μl, 1:10,000 in PBS-TB for 45 min at 4 °C; Sigma). Absorbance was measured at 405 nm (and at 540 nm reference wavelength) using a Tecan Infinite M1000 plate reader after incubation with the substrate disodium 4-nitrophenyl phosphate (100 μl, 3 mM in buffer containing 50 mM NaHCO$_3$ and 50 mM MgCl$_2$ for 60 min at 37 °C; Fluka).

Anisotropy measurement

The assays were performed in black non-binding 96-well plates (Greiner). Then, 2 nM or 10 nM of peptide–sfGFP fusion protein was titrated with increasing concentrations of ArmRP. The concentration of peptide–sfGFP was chosen to be maximally twofold over the respective $K_d$ (CAR.V2) but preferentially below $K_d$ (CAR.V1/V3/V4 and...
CAR2). For the variants, a dilution series of 24 concentrations of dArmRP were used, and fluorescence anisotropy was measured on a Safire II plate reader (Tecan). Data were averaged from four samples, and the anisotropy value from the lowest ArmRP concentration was subtracted for normalization. Data were fitted by a simple one-to-one binding model using SigmaPlot®, using Eq. (4):

\[
[AB] = \left( -\frac{1}{2} \right) \cdot \left( -A_{\text{tot}} + [A_{\text{tot}}] + [B_{\text{tot}}] \right) + \sqrt{\left( A_{\text{tot}} + [A_{\text{tot}}] + [B_{\text{tot}}] \right) ^2 - 4 \cdot [A_{\text{tot}}] \cdot [B_{\text{tot}}]}
\]

where \([AB]\) is the complex formed, \([A_{\text{tot}}]\) and \([B_{\text{tot}}]\) are the concentrations of peptide and ArmRP, respectively, and \(K_d\) is the dissociation constant. \(A_{\text{tot}}\) is the measured anisotropy, \(P_{\text{min}}\) is its minimum in the absence of ArmRP, and \(P_{\text{max}}\) is the value when the peptide is fully bound by protein.

**PDB entry codes**

Coordinates and structure factors have been deposited in the PDB under the following entry codes: 4D4E (YIII(Dq)4CPAF) and 4D49 (YIII(Dq.V1)4CPAF).

**Acknowledgments**

We would like to thank Céline Stutz-Ducommun and Beat Blattmann from the high-throughput crystallization center and the staff from beamlines X06SA and X06DA of the Swiss Light Source for skillful technical support. We would also like to thank Dr. Johannes Schilling for helpful discussions. S.J.F. was supported by the Human Frontier Science Program. This work was financially supported by a Swiss National Science Foundation grant (Sinergia S-41105-06-01).

**Appendix A. Supplementary Data**

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.jmb.2016.09.012.

**References**


ArmRPs for Modular Peptide Recognition


