Computationally designed peptides for self-assembly of nanostructured lattices

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Folded peptides present complex exterior surfaces specified by their amino acid sequences, and the control of these surfaces offers high-precision routes to self-assembling materials. The complexity of peptide structure and the subtlety of noncovalent interactions make the design of predetermined nanostructures difficult. Computational methods can facilitate this design and are used here to determine 29-residue peptides that form tetrahelical bundles that, in turn, serve as building blocks for lattice-forming materials. Four distinct assemblies were engineered. Peptide bundle exterior amino acids were designed in the context of three different interbundle lattices in addition to one design to produce bundles isolated in solution. Solution assembly produced three different types of lattice-forming materials that exhibited varying degrees of agreement with the chosen lattices used in the design of each sequence. Transmission electron microscopy revealed the nanostructure of the sheetlike nanomaterials. In contrast, the peptide sequence designed to form isolated, soluble, tetrameric bundles remained dispersed and did not form any higher-order assembled nanostructure. Small-angle neutron scattering confirmed the formation of soluble bundles with the designed size. In the lattice-forming nanostructures, the solution assembly process is robust with respect to variation of solution conditions (pH and temperature) and covalent modification of the computationally designed peptides. Solution conditions can be used to control micrometer-scale morphology of the assemblies. The findings illustrate that, with careful control of molecular structure and solution conditions, a single peptide motif can be versatile enough to yield a wide range of self-assembled lattice morphologies across many length scales (1 to 1000 nm).

INTRODUCTION

Self-assembly of designed molecules in solution provides marked potential for efficiently achieving complex, robust materials with nanometer precision. Traditional nanomaterial assembly strategies have used small molecules (1, 2) or polymeric (3–5) amphiphiles. Recently developed assembly methods can produce complexity in structure and composition through chemical variation of the assembling molecules (6–9) or the use of hierarchical solution assembly protocols (10–13). Biopolymers offer unique capabilities to encode both local molecular building block structure and long-range material morphology via the design of specific sequences; this design has been applied to DNA (14, 15), polypeptides (16–22), and polymer-biomolecule hybrids (14, 23). Solution assembly of peptides can readily produce “one-dimensional” nanostructures such as fibrils (24–26) and tubes (27–29). New peptide nanostructure formation strategies have used nonnatural peptide sequences (19) and biomimetic strategies using modified natural proteins (30, 31). Much of the peptide work involves the synthesis of new systems and subsequent characterization of the structures they form. However, the a priori design of proteins and peptides that form targeted assemblies is subtle because of the complexities and subtleties of folding and protein-protein interactions. Moreover, these assemblies can be highly sensitive to sequence and mutation. These difficulties have motivated the experimental use of interactions that are more easily programmed at protein interfaces, such as metal coordination (32), to drive intermolecular assembly.

Theoretical and computational methods provide a way to approach the design of intermolecular, noncovalent interactions between self-assembling peptides or proteins in solution to produce materials with predetermined morphologies, including desired point and space-group symmetries (33–36). Nearly all these efforts in assembly design have used variants of natural proteins as building blocks, and different tertiary and quaternary structures are often used for different local geometries in the assembly. Here, we present the computational de novo design of peptides that are robust, easily synthesized, and versatile.

Our aim is to explore the extent to which the information required for folding and intermolecular long-range order can be designed de novo into short peptide sequences, as opposed to the redesign of large natural proteins. The effort is focused on (i) the de novo design of a homotetrameric helix bundle motif that is robust with respect to variation of exterior residues; (ii) the design of the exterior residues to guide the solution assembly of variants of this motif into distinct lattices having rectangular, square, or hexagonal local symmetries; (iii) the experimental characterization and determination of the extent to which the nanostructures are robust with respect to solution-phase conditions; and (iv) the exploration of how solution conditions can be used to control micrometer-scale morphology.

RESULTS

The designed building block motif consisted of a helical homotetramer of four 29-residue peptides arranged with D2 symmetry (Fig. 1). Although
many similar oligomeric helical proteins have been designed and investigated, we seek a structure that is robust with respect to variation of exterior residues, and we opt to design the structure and the hydrophobic core de novo. Candidate bundles were generated via a multiparameter mathematical model of helical coiled coils, with the final bundle structure specified by a set of five defining geometric parameters (see the Supplementary Materials). For each candidate bundle motif, a probabilistic approach was applied to calculate the site-specific probabilities of the amino acids at variable residues (33). The calculations also yield an average energy, \( E \), over sequence probabilities for a given bundle structure (33). \( E \) was used as an objective function in a Monte Carlo search over helix bundle parameters. A helical peptide structure and 11 interior hydrophobic residues were specified (highlighted in gray in Table 1), providing the tetrameric helix bundle motif, or building block, for subsequent design of the material assemblies. The remaining 18 residues were designed in the context of four predetermined material nanostructures: an isolated, water-soluble helix bundle not expected to self-associate (Fig. 1A) and three material assemblies derived from P622, P422, and P222 space-group symmetries (Fig. 1, B to D). Each of these layered space-group symmetries contains \( D_2 \) symmetric positions on which the individual peptide bundles were positioned (Fig. 1, B to D). Calculations were performed using only a single, isolated layer from the corresponding space group. For a given nanostructure symmetry, the variation of the unit cell parameters produced a set of possible lattice structures consistent with the chosen symmetry; the amino acid probabilities and \( E \) were calculated for each assembly structure. From the resulting energy landscape for each type of assembly, energy minima were identified. Within these minima, sequences were identified, where the amino acid with the largest calculated probability was selected at each variable residue position.

In solution, all but one of the computationally designed peptides exhibited the intended assembly properties: the peptide designed using the isolated tetramer template (Fig. 1A) remained soluble and did not form a higher-order structure, whereas those designed in the context of a lattice (Fig. 1, B to D) formed some sort of regular array. Nine
designed sequences were selected for experimental investigation (Table 1). The single sequence designed to form isolated bundles (BNDL_1) formed soluble homotetramers that did not undergo further interbundle assembly. Seven molecules computationally designed to form material assemblies with specific, interbundle packing produced sheetlike nanostructures. No assembly was observed for the sequence P222_4 (Table 1).

Figure 2 presents representative data from four different peptide molecules theoretically designed to produce the four chosen nanostructures in Fig. 1. Figure 2A shows small-angle neutron scattering (SANS) results for BNDL_1 modeled as a short cylinder with dimensions consistent with the designed, tetrameric coiled coil. Analytical ultracentrifugation (AUC) supports the presence of tetrameric helix bundles coexisting with a minority of monomeric peptide (fig. S14). BNDL_1 solutions were experimentally monitored for 3 months, and no interbundle lattice structure was observed. Figure 2B shows the nanostructure formed from assembled P622_6 molecules, which is consistent with hexagonal local symmetry and the targeted unit cell parameters (Table 2). The structure is stabilized by two unique bundle-bundle interfaces that arise from the design around the threefold symmetry axis (Fig. 1B). Both interfaces, one between two antiparallel helices and the other between two parallel helices, are populated by salt bridges, hydrogen bonds, and hydrophobic interactions in the computational design. Figure 2C shows the materials formed from the self-assembly of P422_1 that have the targeted fourfold-like symmetry. Although the fourfold symmetry is clear in Fig. 2B, the experimentally observed unit cell dimensions are larger than predicted (Table 2). Figure 2D shows the materials formed from the self-assembly of P222_1. The observed morphology consists of regular nanosheets that have the targeted rectangular structure, but differences are observed in symmetry and unit cell spacings from what was theoretically designed (Table 2). Although the targeted lattice parameters are not precisely recovered experimentally in each case, together, the data support the use of computational design to realize a set of homotetrameric helix bundles that have predetermined self-assembly properties (for example, sheet-forming tetrameric peptide bundles versus soluble bundles) and distinct, local ordering that is determined by the exterior surface residues of the helical bundle.

The robustness and versatility of the designed bundle-forming peptides were further explored by changing solvent conditions. Different solution conditions, selected on the basis of the physicochemical properties of the bundles, could be used to manipulate interbundle assembly and alter the resulting superstructural morphology of the nanomaterials.

Table 1. Table of computationally determined peptides for solution assembly. Colored rectangles contain eight candidate sequences that were experimentally characterized. Sequences were theoretically designed to produce tetrahelical bundles. BNDL_1 was designed in the absence of any lattice assembly and is expected to remain soluble (brown). The P222 (orange), P422 (green), and P622 (blue) sequences were designed in the presence of lattices of corresponding symmetry. The remaining P222_9 and P422_1 sequences contain covalently modified termini. P222_4 is the only sequence candidate that did not behave as predicted and could not be assembled into a nanostructure in the solution conditions used for the other peptides. The heptad repeat positions (abcdefg) of all peptides are shown in the table heading. The designed, hydrophobic interior residues of the motif shown are highlighted in gray. MW, molecular weight.

| Name    | abcdefg | abcdefg | abcdefg | abcdefg | a    | pI | MW(dalton) |
|---------|---------|---------|---------|---------|------|----|------------|
| BNDL_1  | DEEIRRM | AEIIRQM | AERIQOM | AEIQIQE | A-NH2| 4.32| 3560       |
| P222_1  | DGKIEGM | AEAIKKM | ANNIEQM | AGWIWGE | A-NH2| 4.79| 3192       |
| P222_9  | DGRIEGM | AEAIKKM | AYNIADM | AGRIWGE | A-NH2| 7.13| 3168       |
| P222_10 | DGKIEGM | ADRIRRM | ARNIEDM | AYiYRE  | A-NH2| 4.89| 3404       |
| P422_1  | DQEIRQM | AEWIKKM | AQMIDKM | AHRHIDR | A-NH2| 9.75| 3572       |
| P622_1  | DEIJKQM | AHWGEM | AGQINKM | ASESIAE | A-NH2| 5.50| 3217       |
| P622_2  | DEIIKLM | AEWIKKM | AGNIDKM | AKIDRDE | A-NH2| 9.59| 3407       |
| P622_6  | DEIIKLM | ADQIKKM | AWIMDRE | AEKIDRE | A-NH2| 7.75| 3517       |
| P222_9_Ac | Ac-DGRIEGM | AEAIKKM | AYNIADM | AGRIWGE | A-NH2| 4.61| 3210       |
| P422_1_Ac | Ac-DQEIRQM | AEWIKKM | AQMIDKM | AHRIDRE | A-NH2| 8.12| 3614       |
| P222_9_2Gly | GG DGRIEGM | AEAIKKM | AYNIADM | AGRIWGE | A-NH2| 7.13| 3282       |
| P222_9_4Gly | GGGG DGRIEGM | AEAIKKM | AYNIADM | AGRIWGE | A-NH2| 7.13| 3396       |
| P222_9_6Gly | GGGGGG DGRIEGM | AEAIKKM | AYNIADM | AGRIWGE | A-NH2| 7.13| 3510       |
| P222_4   | DEIKEM ADQIKRM | ANEIEEM | AGWIWGE | A-NH2 | 4.38| 3422       |
For example, Fig. 3 (A and B) reveals that the micrometer-scale morphology of P622_2 can be manipulated simply by first melting the secondary structure at 80°C and subsequently quenching the solution to two different temperatures. Smaller particles were formed at the higher quenching temperature (50°C) than at the lower quenching temperature (25°C). The data suggest that a higher temperature results in a much slower assembly process. Figure 3 (C and D) reveals the sensitivity of select bundle assemblies to changes in pH. The molecule P222_9 has a theoretical isoelectric point (pI) = 7 and was assembled under two solution conditions that differed only in pH. The assembly at pH 7 resulted in two-dimensional, plate-like growth, whereas the assembly at pH 10 yielded anisotropic growth and the formation of long, needle-like structures. The thickness of the needles prevented clear, high-magnification lattice imaging, but the layer spacings of the underlying lattice (fig. S15) were consistent with the nanostructure observed for the molecule P222_9 (Figs. 1 and 2). The results suggest that, at pH 10, there is a clear preference in growth direction during helix bundle solution assembly.

Covalent alteration of the original designed peptides can also be used to probe the robustness of the assembly and to modulate assembly at particular solution conditions. For example, acetylation of the N terminus of P222_9 (denoted P222_9_Ac; Table 1) reduces its theoretical pI from pI = 7 (P222_9) to pI = 4 (P222_9_Ac), allowing for assembly at low pH. At pH 4.5, P222_9 remains dissolved, whereas P222_9_Ac (Fig. 3E) assembles into the same nanostructure as that observed for P222_9 at pH 7 (fig. S16). Similarly, P422_1_Ac forms plate nanostructures at pH 8 (Fig. 3F), whereas P422_1 assembles at pH 10 (Fig. 2B). Therefore, one can use covalent modifications of the designed peptide sequences to alter solution conditions in which nanostructures can be formed. Adding residues to the termini of the originally predicted sequences did not disrupt solution self-assembly. Specifically, sequences

Table 2. Lattice parameters of the self-assembling peptides from the design in comparison with those determined from analysis of Fourier transforms of the TEM images in Fig. 2. a and b denote the dimensions of the two-dimensional unit cell, and \( \gamma \) denotes the interior angle defined by sides a and b.

| Design | TEM | 
|--------|-----|
|        | a (nm) | b (nm) | \( \gamma \) (°) | a (nm) | b (nm) | \( \gamma \) (°) |
| P622_6 | 4.57   | 4.57   | 120              | 4.5 ± 0.3 | 4.5 ± 0.3 | 112.7 ± 0.4 |
| P422_1 | 3.12   | 3.12   | 90               | 4.2 ± 0.2 | 3.9 ± 0.2 | 88.9 ± 0.9  |
| P222_1 | 2.09   | 2.00   | 90               | 3.3 ± 0.3 | 3.2 ± 0.3 | 100.4 ± 0.9 |

For example, Fig. 3 (A and B) reveals that the micrometer-scale morphology of P622_2 can be manipulated simply by first melting the secondary structure at 80°C and subsequently quenching the solution to two different temperatures. Smaller particles were formed at the higher quenching temperature (50°C) than at the lower quenching temperature (25°C). The data suggest that a higher temperature results in a much slower assembly process. Figure 3 (C and D) reveals the sensitivity of select bundle assemblies to changes in pH. The molecule P222_9 has a theoretical isoelectric point (pI) = 7 and was assembled under two solution conditions that differed only in pH. The assembly at pH 7 resulted in two-dimensional, plate-like growth, whereas the assembly at pH 10 yielded anisotropic growth and the formation of long, needle-like structures. The thickness of the needles prevented clear, high-magnification lattice imaging, but the layer spacings of the underlying lattice (fig. S15) were consistent with the nanostructure observed for the molecule P222_9 (Figs. 1 and 2). The results suggest that, at pH 10, there is a clear preference in growth direction during helix bundle solution assembly.
of two, four, and six glycine residues were added to the N terminus of P222_9 (Table 1), resulting in quite uniform assemblies (Fig. 3, G and H) with similar lattice structures to those observed for the unmodified P222_9 (Fig. S17). These observations reveal the robust nature of these theoretically designed peptides and their material assemblies. Together, the data from Fig. 3 exemplify that multiple modifications of the theoretically predicted sequences remain competitive for forming the selected nanostructures.

DISCUSSION

The data from Figs. 2 and 3 and the Supplementary Materials support the fact that theoretically designed sequences listed in Table 1 self-assemble to form predetermined sheet nanostructures or soluble bundles. Specifically, the sequence BNDL_1, targeted to form soluble, nonassociating bundles, formed homotetramers in solution but did not exhibit further lattice assembly. Similarly, the sequence P622_6 was designed to form two-dimensional sheets with hexagonal local symmetry and formed a nanostructure consistent with that identified in the computational design of the sequences; this consistency is evidenced in the similarity of the unit cell parameters of the theoretical model to those derived from analysis of TEM data. As shown in Fig. 2, the molecules designed to produce two-dimensional sheets with P422 symmetry assemble into a two-dimensional sheet nanomaterial with local fourfold bundle packing symmetry. However, the lattice spacing formed experimentally is larger than that expected from the computational design. Similarly, the molecules predicted to yield the two-dimensional materials with P222 symmetry assembled into sheet nanostructures. However, the lattice symmetry and unit cell parameters were different from those identified in the computational modeling. Additional calculations using the P422 molecule sequence within a related fourfold symmetry, P4, reveal a local energy minimum at the experimentally determined unit cell parameters (Fig. S18). The distance between the C2 symmetry axes of neighboring bundles within each model structure is essentially indistinguishable: 2.2 nm (P4) versus 2.2 nm (P422). With the P4 structure, the unit cell contains four helical bundles (16 peptides) as opposed to two helical bundles (8 peptides) in the original model with P422 symmetry, which is consistent with the larger unit cell dimensions observed experimentally. Similar calculations involving P222 within a related twofold symmetry, P2, reveal a local minimum that is consistent with the experimental observations (Fig. S19). More effort is needed to refine design methods, recognize deterministic molecular attributes, and develop assembly pathways that will reproducibly funnel the molecules into predetermined nanostructures during solution assembly. Nonetheless, the designed assemblies are robust over a wide range of pH and temperature values and are resilient with respect to chemical modification at the peptide termini. Thus, the control of solution conditions (pH and temperature), primary sequence, and chemical modification of peptides can be used to control solution-phase assembly and the final material morphology.

We have presented an approach for the de novo design of peptide assemblies that assemble into nanomaterials with predetermined local structures. We computationally designed a versatile, helical, homotetrameric building block structure that was stabilized largely by the formation of a hydrophobic interior. Subsequent computational determination of the bundle exterior residues resulted in solution-assembled materials with predetermined morphologies (two-dimensional sheet versus soluble bundle) and distinct, local nanostructures that varied with the bundle exterior sequence. Integrated theoretical and experimental studies will guide future refinements for predictably achieving specific, desired nanostructures and elucidating the principles underlying their formation.
In addition to achieving symmetric assemblies of biopolymers, the display of a wide variety of chemical functional groups within designed assemblies offers abundant opportunities for hierarchical pathways for nanomaterial production, such as biopolymer-templated growth and/or assembly of inorganic phases with nanoscale precision. Computational design combined with the experimental control of assembly pathways has the potential to provide exquisite control over new materials with desired nanostructures.

MATERIALS AND METHODS

Peptide synthesis
Peptides were prepared at a 0.25-mmol scale on a Rink amide resin using an AAPPTec Focus XC synthesizer (AAPPTec). Standard Fmoc-based protocols were used. Peptides were deprotected for 5 min and then for 10 min with 20% piperidine in dimethylformamide (DMF). The coupling reaction was conducted for 40 min with 4 eq of the appropriate amino acid dissolved in N-methyl-2-pyrrolidone (NMP) (5 ml), 3.8 eq of O-(6-chlorobenzotriazol-1-yl)-N,N,N′,N′-tetramethyluronium hexafluorophosphate (HCTU) dissolved in DMF (2.5 ml), and 8 eq of disopropylethylamine dissolved in NMP (1 ml). Five washes were performed in between steps with 50:50 (v/v) DMF/methylene chloride (12 ml) for the first two washes and with DMF (10 ml) for the last three. Amino acids, resins, and activators were purchased from ChemPep and were used as received. All solvents were of analytical grade (Fisher Scientific). Peptide cleavage was achieved by shaking the peptide solutions for 2 hours in a cleavage cocktail comprising (by volume) 95% trifluoroacetic acid (TFA), 2.5% triisopropylsilane, and 2.5% Milli-Q water. The peptide was then precipitated by adding the cleavage cocktail and cleaved peptide to diethyl ether. The mixture was then centrifuged, and the supernatant was discarded. The process of suspending in diethyl ether, centrifuging, and discarding the supernatant was repeated a total of three times. The resulting peptide was then dissolved in water and lyophilized.

Peptide purification
Purification was performed via reversed-phase high-performance liquid chromatography (HPLC) using a BEH130 Prep C18 10-μm column (XBridge, Waters Corporation). Crude peptides were dissolved in Milli-Q water containing 0.1% (v/v) TFA and were filtered (0.20-μm filter, Corning Inc.) before HPLC injection. The products were subjected to an elution gradient [Quaternary Gradient Module (Waters 2545), Waters Corporation] of 100% solvent A [0.1% (v/v) TFA] to 30% solvent A within 60 min; solvent B consisted of acetonitrile with 0.1% (v/v) TFA. Fractions were detected using ultraviolet-visible detection at 214 nm (Waters 2489, Waters Corporation) and collected (Waters Fraction Collector III, Waters Corporation). The collected fractions were examined by electrospray ionization–mass spectrometry (LCQ Advantage Mass Spectrometer System, Thermo Finnigan) with an autosampler system (Surveyor Autosampler, Thermo Finnigan). Pure fractions were combined and lyophilized.

Analytical HPLC
Purity analysis was performed via reversed-phase analytical-scale HPLC using a BEH C4 3.5-μm column (XBridge, Waters Corporation). Peptides were dissolved in 80:20 (v/v) Milli-Q water/acetonitrile containing 0.1% (v/v) TFA and filtered with a 0.20-μm filter (Corning Inc.) before injection. The products were subjected to a linear elution gradient (Waters 600 Controller, Waters Corporation) of 80% solvent A [Milli-Q water with 0.1% (v/v) TFA] to 5% solvent A in 70 min; solvent B consisted of acetonitrile with 0.1% (v/v) TFA. Fractions were detected using a photodiode array detector (Waters 2996, Waters Corporation) tuned at 214 nm.

Circular dichroic spectroscopy
Secondary structures and the temperature-dependent behavior of the synthetic peptides were analyzed using circular dichroic spectroscopy on a Jasco J-820 spectropolarimeter (JASCO Inc.). Sample solutions were prepared at 0.1 mM concentration in 10 mM buffer appropriate for the desired pH and were transferred into an absorption cuvette with a path length of 1 mm (110-QS, Hellma Inc.). Pure buffer solutions were used for the background correction. Sample spectra were recorded from 190 to 250 nm at desired temperatures. Data points for the wavelength-dependent circular dichroic spectra were recorded at every nanometer with a bandwidth of 1 nm and a response time of 4 s for each data point. The ellipticity at 222 nm was used to monitor the temperature-dependent unfolding and refolding of the peptides. Data points for the scans were recorded at 222 nm at 1-min intervals. The mean residue ellipticity, [θ]_{MRE} (deg cm² dmol⁻¹), was calculated using the peptide concentration, number of amino acid residues, and cell path length.

Transmission electron microscopy
Carbon-coated 200-mesh copper grids (CF200-Cu, Electron Microscopy Sciences Inc.) were freshly treated by glow discharge using a plasma cleaner (PDC-32G, Harrica Plasma Inc.) before sample preparation. Three microilters of the sample suspension were applied onto the grids. After ~5 min, any remaining liquid was blotted from the edge of the grids using filter paper. Then, 3 μl of Milli-Q water was applied to the grids and blotted immediately to remove excess un assembled peptides and buffer salts. The grids were incubated under ambient conditions for another 30 min before TEM observation (Tecnai 12, FEI) or staining. To negatively stain the grids, 3 μl of an aqueous solution of uranyl acetate (1 weight percent) was applied to the cast-film grids, incubated for 20 to 30 s, and then blotted with filter paper. The stained grids were allowed to sit for at least 10 min under ambient conditions before TEM observation. Positive staining of the samples was achieved by exposing the dried cast-film grids to ruthenium tetroxide aqueous solution vapor [0.5% (w/v)] for 5 to 10 min. The transmission electron microscope was operated at an acceleration voltage of 120 kV, and all images were recorded digitally using a Gatan 791 Multiscan side-mounted charge-coupled device camera. Lattice plane spacings were determined by conducting an FFT of the area of interest of a TEM micrograph using DigitalMicrograph software version 2.3 (Gatan Inc.) and the PASAD plug-in (37). FFT intensity was plotted relative to radial distance from the origin as well as relative to azimuthal angle. Corresponding real space values of interplanar distances and angles were calculated to estimate unit cell parameters. Uncertainties of the unit cell parameters were approximated by the size of the maxima in the FFT. Fourier-filtered images (lower insets in Fig. 2) were obtained by applying masks around Fourier frequencies with distinct intensity maxima within the respective FFT and then performing an inverse FFT.

Small-angle neutron scattering
SANS measurements were conducted at the National Center for Neutron Research [National Institute of Standards and Technology (NIST),
Gaithersburg, MD) on the NG-7 30-m SANS beamline. Samples were dissolved in 10 mM borate buffer (pH 10) prepared in D2O and heated to 80°C for 1 hour to melt any organized secondary structure/ aggregated structure and were subsequently cooled to room temperature for intermolecular assembly. A neutron beam with a mean wavelength of 6 Å was defined using a mechanical velocity selector. The wavelength spread (Δλ/λ) was 0.15 at full width at half maximum. The 640 mm × 640 mm He proportional counter used has a spatial resolution of 5.08 mm × 5.08 mm. Sample-to-detector distances of 1, 4, and 13 m were used to provide a q range of approximately 0.004 to 0.50 Å−1, where q is the scattering wave vector defined by q = (4/λ) sin(θ/2).

Data obtained with this instrument were corrected for background noise and radiation, detector inhomogeneity, and empty cell scattering. Intensities were normalized to an absolute scale relative to the empty beam transmission. The uncertainties of individual data points were calculated statistically from the number of averaged detector counts. See the Supplementary Materials for a description of data fitting and analysis.

**Analytical ultracentrifugation**

AUC experiments were run on a Beckman Coulter ProteomeLab XL-I instrument. Four hundred microliters of 0.5 mM peptide solution in buffer (10 mM borate buffer with extra 50 mM NaCl to screen long-range Coulombic interactions; pH 10) was transferred into a two-channel cell equipped with sapphire windows. The cells were mounted into a four-cell An-60 Ti analytical rotor and equilibrated at 20°C for 2 hours in the rotor chamber. Sedimentation velocity experiments were carried out at 50,000 rpm at 20°C using 800 scans with 1 scan per minute per sample. The changes of the concentration profiles along the cell radius were monitored using Rayleigh interference optics. All data were analyzed using the SEDFIT package version 14.81 (38).

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/2/9/e1600307/DC1

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RESEARCH ARTICLE

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