Computational design and experimental verification of a symmetric protein homodimer

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Homodimers are the most common type of protein assembly in nature and have distinct features compared with heterodimers and higher order oligomers. Understanding homodimer interactions at the atomic level is critical both for elucidating their biological mechanisms of action and for accurate modeling of complexes of unknown structure. Computation-based design of novel protein–protein interfaces can serve as a bottom-up method to further our understanding of protein interactions. Previous studies have demonstrated that the de novo design of homodimers can be achieved to atomic-level accuracy by β-strand assembly or through metal-mediated interactions. Here, we report the design and experimental characterization of α-helix–mediated homodimer with C2 symmetry based on a monomeric Droso phila engrailed homeodomain scaffold. A solution NMR structure shows that the homodimer exhibits parallel helical packing similar to the design model. Because the mutations leading to dimer formation resulted in poor thermostability of the system, design success was facilitated by the introduction of independent thermostabilizing mutations into the scaffold. This two-step design approach, function and stabilization, is likely to be generally applicable, especially if the desired scaffold is of low thermostability.


Computational design tools use a bottom-up approach that allows for the testing of hypotheses on the relationships between amino acid sequence, protein structure, and biological function. Here, we exploited two computational methods, protein docking and protein sequence optimization, to create a favorable protein–protein interaction between two identical proteins, resulting in a novel homodimer. A stepwise approach proved useful: scaffold stabilization followed by interface design to achieve homodimerization. Our results suggest that for some proteins, stabilization may be required for the successful design of functionality.


The authors declare no conflict of interest.

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Data deposition: The atomic coordinates have been deposited in the Protein Data Bank, www.pdb.org (PDB ID codes 2MG4 and 4NDL).

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Significance

Protein–protein interactions play a central role in nearly all biological processes, including cell signaling, immune responses, regulation of transcription and translation, and cell–cell adhesion. Improving our understanding of protein–protein interactions is therefore an important component to advancements in both basic research and applications in the pharmaceutical, chemical, and biotechnology industries. The increasing availability of high-resolution structures has led to the identification of unique features of protein–protein interactions (1–3). Specific interfacial residues that contribute to most of the binding energy (“hot spots”), networks of hydrogen bonds, and shape complementarity have all been identified as important. These features have therefore been incorporated into many protein docking and protein design algorithms (4, 5). Protein docking algorithms have been used successfully to screen millions of docking positions and to identify the correct (near-native) structures (6). Computational design tools have also exploited our knowledge of protein–protein interactions to design enhanced affinity or altered specificity successfully (7, 8), to graft binding motifs onto a design of an α-helix–mediated C2 symmetric homodimer based on a monomeric globular protein scaffold.

Computational protein design tools use a bottom-up approach that allows for the testing of hypotheses on the relationships between amino acid sequence, protein structure, and biological function. Here, we exploited two computational methods, protein docking and protein sequence optimization, to create a favorable protein–protein interaction between two identical proteins, resulting in a novel homodimer. A stepwise approach proved useful: scaffold stabilization followed by interface design to achieve homodimerization. Our results suggest that for some proteins, stabilization may be required for the successful design of functionality.


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protein–protein interactions. Although there are empirical rules for archetypal coiled-coil oligomerization (27), a general sequence–structure relationship that could be applied to an arbitrary scaffold has not yet been found. Furthermore, as shown by Keating and coworkers (28), predicting parallel or antiparallel helix-helix homodimers using computational modeling is challenging.

The similarity between parallel and antiparallel helix-helix structures and the high hydrophobicity of homodimers make it difficult to distinguish between the different conformational states, particularly if they are strongly competing with each other and only one of the states is explicitly designed. For example, Karamicolas et al. (29) computationally designed a novel protein–protein interface with tightly packed hydrophobic residues. The crystal structure, however, revealed that the orientation of one of the partners was rotated almost 180° relative to its position in the design model. These results underscore the difficulty of excluding unwanted competing states in the design of protein–protein interactions.

Here, we design a C2-symmetric homodimer from a helical monomeric protein, Drosophila melanogaster engrailed homeodomain (ENH). This small helix-turn–helix protein domain binds a specific sequence of dsDNA (30) and has been used as a model for theoretical and computational studies (31–33). Computational protein design (CPD) endeavors often begin with a thermostabilized scaffold because designs for targeted function (e.g., catalytic activity, ligand–protein binding affinity) have been shown to decrease protein stability significantly (34). Poor stability, in turn, often results in aggregation and can be problematic for protein expression and/or experimental characterization. The 51-aa fragment of WT ENH used in this study has a melting temperature (Tm) of 49 °C (35), which is low for a design scaffold. Indeed, we found that our initial attempt to build a homodimer using this scaffold resulted in a protein, ENH-c2a, that expresses in inclusion bodies even at temperatures as low as 16 °C. For subsequent designs, we used the thermostabilized variant ENH NC3-NCap, a computationally designed protein that has a Tm of 89 °C (35). We applied a symmetric docking program based on a fast Fourier transform (FFT) algorithm (36) and designed the interface between four helices (two from each molecule) so that they would associate as a four-helix bundle. The final design (ENH-c2b) was experimentally characterized and shown to be a monodisperse homodimer with a Kd of ∼130 nM. The solution NMR structure reveals that the helical interface exhibits parallel packing consistent with the design model.

Results

Scaffold Selection. Due to our incomplete understanding of structure–function relationships and limitations in our ability to model proteins accurately, the success rate for CPD is generally rather low. This low success rate is particularly true for the de novo design of functional proteins, such as enzymes or proteins designed to interact with a specific target (37). Many de novo design studies therefore use a comprehensive approach that screens in silico scaffolds in the PDB, often resulting in tens to hundreds of candidate scaffolds (12, 16, 38, 39). These candidates are then used as the basis for computational design, with the top designs screened experimentally (e.g., for target binding affinity or catalytic activity). This approach has the advantage of providing a large amount of data that can be used to test design protocols (22, 40). Alternatively, focusing on a single scaffold allows the design to take specific features of the scaffold into account. For example, Privett et al. (34) used an iterative approach on a single scaffold that resulted in the de novo design of a Kemp eliminase that was further optimized by directed evolution to yield an enzyme with native-like kinetic parameters (41). ENH was selected as the target scaffold for this study because of its ability to bind to DNA, which could be exploited in future designs to create protein-DNA nanomaterials that self-assemble through noncovalent interactions. ENH binds DNA primarily with its third helix, leaving the surfaces of the first two helices available for design (30).

Design Protocol. Fig. 1 shows the steps we used to design and characterize a C2-symmetric homodimer. After selecting the scaffold protein, the surface side chains were pruned to CB and the atomic radii were parameterized based on known C2-symmetry homodimers. We then applied a docking procedure, constrained for C2 symmetry, to generate initial models based on an FFT algorithm (36). FFT docking allowed for the efficient search of the translational and rotational 6 df. Approximately 1010 models were screened and ranked by shape complementarity. The top 200 candidates were clustered into 11 groups according to the pairwise rmsd values of the structures. Finally, these clusters were visually inspected, and one model was chosen for homodimer design. In the selected homodimer model, the first helix of each monomer pairs to form a parallel packing arrangement with the helices separated by 10 Å, similar to the 9.8-Å separation found in naturally occurring coiled-coil dimers (42) even though the design process did not explicitly consider the heptad repeat rules that govern coiled-coil interactions (24). Taken together, the first two helices of each monomer form a four-helix bundle structure (Fig. 1).

Symmetric sequence optimizations were applied to the 22 interfacial residues (11 on each side) of the homodimer model. To recapitulate the balance between nonpolar (∼65%) and polar (∼35%) amino acids seen in natural homodimers (23), the value of the force-field parameter most responsible for limiting the burial of polar surface area, σp, was doubled relative to its standard setting (43).

Initial homodimer design was performed using WT ENH as the starting scaffold, resulting in a protein, ENH-c2a (Table 1), that could only be expressed in inclusion bodies (Fig. S1). Second-generation designs were performed using a previously designed thermostabilized variant of ENH, NC3-NCap, for the scaffold (35). A computationally designed library consisting of 128 variants of NC3-NCap was generated (Table S1), fused to YFP, and screened using a homo-FRET assay (Fig. S2A). Two variants resulting from the screen were characterized via size exclusion chromatography and analytical ultracentrifugation sedimentation velocity experiments; one proved to be a dimer, and the other was a tetramer (Fig. S2 B and C). In a related paper (44), we
showed that the dimer, ENH_DsD, is domain-swapped and that it could be converted to a non-domain-swapped dimer through the introduction of proline at residue 23. The resulting sequence, ENH-c2b (Table 1), expressed well in the soluble fraction.

Biophysical Characterization of ENH-c2b. The designed proteins were characterized for soluble expression, secondary structure, thermostability, and oligomeric state. SDS/PAGE gels of purified ENH, ENH-c2a, and ENH-c2b are shown in Fig. S1. Although ENH could be expressed in the soluble fraction at 16 °C, ENH-c2a showed no soluble expression under the same conditions. In contrast, ENH-c2b expressed well at 37 °C, with yields of over 5 mg/L culture. CD spectroscopy revealed that ENH-c2b is helical, folds reversibly, and has a Tm of ~62 °C (Fig. 2). Introduction of the designed interface reduced the Tm of the scaffold by ~26 °C (Table 1).

Analytical ultracentrifugation sedimentation velocity experiments showed that ENH-c2b is a monodisperse dimer at a concentration of 5 μM (Fig. 3A). The Kd for dimer formation was determined using a tryptophan fluorescence-based homo-FRET assay and was found to be 129 ± 64 mM (Fig. 3B), which is similar to the Kd values reported for other de novo-designed protein interfaces (before affinity maturation) (12, 13, 16).

Structure Determination of ENH-c2b. X-ray crystallography attempts using an ENH-c2b variant with an extended and Histagged N terminus derived from the expression vector (pET28a) resulted in a dimer structure dominated by crystal packing not matching the design target and an interface area in the design region of only 699 Å2 (Fig. S3 and Table S2). Heteronuclear single quantum coherence (HSQC) NMR of 13C and 15N uniformly labeled, freshly prepared ENH-c2b showed a well-folded protein with sharp peaks that broadened over time. Adding six WT ENH amino acids (MEKRPR) at the N terminus, a Gly, and an eight-residue Strep-tag II at the C terminus of ENH-c2b (ENH-c2b-Strep) greatly enhanced its long-term stability. All NMR spectra showed only one set of chemical shifts for all residues, reflecting the underlying symmetry of ENH-c2b-Strep. Unambiguous chemical shift assignments were obtained for most of the backbone nuclei; however, peaks were missing for the last three residues, 21–23 (FYF), at the end of helix-1. This finding is consistent with the fact that the chemical shifts of aromatic residues are highly sensitive to their side-chain conformations and can be easily broadened if multiple conformations exchange on the NMR time scale (45).

We determined the structure of the ENH-c2b-Strep homodimer using ψ/φ angle, hydrogen-bond, NOE, and C2-symmetry restraints (Table S3). Final coordinates of 10 models were deposited in the PDB with ID code 2MG4 (Fig. 4). The overall completeness of all (intramolecular and intermolecular) NOE restraints is 57%, 37%, and 22% at 3-Å, 4-Å, and 5-Å cutoff distances, respectively (Table S4). For comparison, the average completeness for the 97 entries reported by Doreleijers et al. (46) is 68 ± 14%, 48 ± 13%, and 26 ± 9%, at 3-Å, 4-Å, and 5-Å cutoff distances, respectively. There are 26 and 33 unambiguous intermolecular NOE restraints that have zero or one violation, respectively, in the 10 models of the NMR ensemble (Table S5). The intermolecular completeness of NOE restraints is 52% at a 4-Å cutoff distance (Table S6) compared with the average value of 37% reported by Doreleijers et al. (46).

Each monomeric subunit superimposes with WT ENH with an overall backbone rmsd of 2.5 Å, with the greatest deviation for the loop between helix-1 and helix-2, residues 21–23 (at the end of helix-1), and the N terminus (Fig. 5A and B), which corresponds to regions lacking restraints and that show the greatest structural variation in the NMR ensemble (Fig. 4). The helices comprising the dimer interface, helix-1 and helix-2, align well with the design model (Fig. 5A), whereas the orientation of helix-3 deviates from the orientation of the model (Fig. 5B). Compared with ENH, the most structurally defined regions of ENH-c2b-Strep (helix-1, helix-2, helix-3, and the loop between helix-2 and helix-3) have a backbone rmsd of 1.4 Å. The solution structure of the ENH-c2b-Strep dimer shows parallel helix-helix packing between helix-1 of each subunit, as in the homodimer design model. Compared with the model, the backbone rmsd of the most structurally defined regions (as discussed above) is 2.2 Å (Fig. 5C and D). The axial orientations of the four helices (helix-1 and helix-2 from each subunit) are nearly identical with those helices in the model (Fig. 5C). The interface area is 2,189 Å2, which falls in the range of natural single-patch homodimers (2,740 ± 1,240 Å2) (23). Nonpolar residues constitute 62% of the interface, close to the average value of 65% for natural homodimers (23). A NOESY experiment designed to retain only intermolecular interactions revealed several nonpolar interfacial residues that are likely to be important for dimerization, including Ala16, Leu19, Ala20, and Leu39 (Fig. 6). Structural alignment of only one of the subunits of the dimer emphasizes the differences between the structure of ENH-c2b-Strep and the design model (Fig. 5E and F).

Discussion

Stranges and Kuhlman (21) recently reviewed the computational design of novel protein–protein interfaces and pointed out the
challenges this burgeoning field faces (21). Of 147 protein–protein interaction designs, only four were confirmed successful by X-ray crystallography (i.e., the solved structure matched the design model). All of the successful designs shared a number of common features: They exhibited fewer polar atoms (<40%) and fewer buried hydrogen bonds at the designed interface than those polar atoms and buried hydrogen bonds seen in the failed designs. This reduced number of buried hydrogen bonds in the successful designs is in contrast to what is typically observed in natural dimers. Our homodimer design had similar characteristics: 62% nonpolar atoms and no buried H-bonds at the interface. This higher interfacial hydrophobicity is expected, given that three of the five successful cases (including the one reported here) were designs for homodimers, which naturally exhibit high nonpolar content at the interface (22). The other two successful cases (both heterodimer designs) also exhibited high interfacial hydrophobicity (73% and 74%), which is likely related to the natural hydrophobicity of the targeted binding surface, the stem region of influenza HA (12, 15). However, designing largely hydrophobic binding surfaces can lead to failure in achieving the targeted structure. For example, the designed protein could bind to the target protein in the wrong orientation (29) or bind to itself to form undesired oligomers (12). In a case described in a related paper (44), when fused with YFP at the C terminus, ENH-c2b forms an unexpected but well-defined homodimer with dimerization interfaces formed between the ENH fragment and YFP. In addition, ENH-c2b with an extended N-terminal tag forms a dimer with an interface that appears to be dominated by crystal packing forces (as described above). Nonetheless, successful designs of protein–protein interfaces based largely on hydrophobic interactions do exist, whereas the successful design of largely polar interfaces has yet to be fully demonstrated. Recently, Procko et al. (47) designed a protein inhibitor that binds to a hydrophilic patch on lysozyme with high affinity; however, this complex has yet to be structurally validated.

An interesting feature of the five successful designs is that the designed interfaces mainly involve the association of well-defined elements of secondary structures. Both of the successful HA heterodimer designs described above have “hot spots” on their helices that bind to helical structures on the target HA stem. Of the three successful homodimer designs to date, one is between two helices (this study), one involves metal–protein interactions mediated by helices (13), and one uses two exposed β-strands to form the homodimer (16). Loops can also be exploited in protein–protein interactions, as demonstrated by the widespread use of loops in antibody–antigen interactions. Many computational loop designs have been attempted; however, thus far, none have resulted in dimerization (12). In a community-wide assessment of protein–protein designs, Haliloglu and coworkers (22) found that many of the failed designs contain more loops and turns than successful designs, and that the higher flexibility of loops makes adopting a particular designed conformation difficult. Noninterfacial loop designs have demonstrated success (48–50). However, the design of interfaces involving loops appears to be more challenging, because the recognition-induced conformational changes that loops can undergo upon association with another protein are still poorly understood, and hence poorly modeled.

A significant difference between the design reported here and the four successful designs reported previously is that the accuracy of our design, with an rmsd of 2.2 Å, is lower than the 1.0- to 1.80-Å rmsd range reported for the other designs. The success of the four more accurate designs may be due to their incorporation of specific hot-spot residues or anchoring interactions that steered the formation of a high-affinity dimer. For example, the two HA heterodimer designs used predefined hot-spot residues to match the specified locations on the target patch (12) and the homodimer designs used either β-strand-mediated hydrogen bonds (16) or metal chelation to anchor the homodimerization interface (13). All of these designs exploited very specific pairwise interactions to facilitate and guide complex formation, which could be a general model that will be useful for future work.

As noted above, our initial design attempts using the WT ENH sequence as the parent for incorporating CPD-directed mutations resulted in a variant, ENH-c2a, that was not solubly expressed even at reduced temperatures, presumably because the introduced mutations compromised the stability of the protein. Introduction of a scaffold stabilization step into the design process (Fig. 1) led to the use of NC3-Ncap, a previously reported stabilized variant of ENH (35) that was able to support the mutations designed to confer homodimerization successfully. Thus, at least in this case, scaffold stabilization facilitated successful functional design. CPD has proved to be useful for designing proteins with improved stability (e.g., 51, 52). Designing for function, such as binding or catalytic activity, however, can lead to significant protein destabilization (34). In one example, Fleishman and coworkers (5) designed 88 proteins to bind to HA and found that 50% of them could not be solubly expressed in Escherichia coli. In the absence of highly stabilized scaffolds,

Fig. 3. Characterization of ENH-c2b oligomeric. (A) Sedimentation velocity experiment at 5 μM with curve fit using the c(M) (continuous distribution of molecular mass) model. (B) Tryptophan homo-FRET assay: circles, experimental data; dashed line, curve fit obtained using a monomer-dimer equilibrium model.

Fig. 4. Solution NMR structure ensemble showing the 10 lowest energy models for the core 51 amino acids (PDB ID code 2MG4). C, C terminus.
All proteins were expressed using deleted and atomic radii of the remaining atoms adjusted length scans and temperature de-CD studies were performed on an Aviv 62A DS spec-ENH-c2b was analyzed on an XL-1 analytical C NMR spectrum showing intermolecular NOE restraints obtained by a ∼ www.pnas.org/cgi/doi/10.1073/pnas.1505072112 ∼ 13 Comparison of ENH-c2b averaged minimized solution NMR structure of 47% compared with WT ENH. fragment used as the design scaffold and has a sequence identity design, ENH-c2b, is 13 °C more stable than the WT ENH was first stabilized and subsequently homodimerized. The final design, ENH-c2b, is a monodisperse dimer with a C2-symmetry homodimer via helical interactions. The successful design model with the protein interface forming a four-helix bundle. The homodimer design was achieved using a two-step computational approach in which the WT protein, ENH, was first stabilized and subsequently homodimerized. The final design, ENH-c2b, is 13 °C more stable than the WT ENH fragment used as the design scaffold and has a sequence identity of 47% compared with WT ENH.

Conclusions

This work represents an early example of the de novo design of a C2-symmetry homodimer via helical interactions. The successful design, ENH-c2b, is a monodisperse dimer with a Kd of ~130 nM. The solution NMR structure is generally consistent with the design model with the protein-protein interface forming a four-helix bundle. The homodimer design was achieved using a two-step computational approach in which the WT protein, ENH, was first stabilized and subsequently homodimerized. The final design, ENH-c2b, is 13 °C more stable than the WT ENH fragment used as the design scaffold and has a sequence identity of 47% compared with WT ENH.

Materials and Methods

Protein Docking and Computational Design. The ENH crystal structure (PDB ID code 1ENH) was used as the scaffold for homodimerization with side-chain atoms beyond Cb deleted and atomic radii of the remaining atoms adjusted as follows: N, 1.4 Å; O, 1.3 Å; C, 1.75 Å; Ca, 2.35 Å; and Ca, 2.15 Å. A symmetric docking program based on an FFT algorithm was applied. One high-scoring model was selected for computational designs. ORBIT CPD software was used for stability designs for both ENH and NC3-NCap. Initial interface designs were also done using ORBIT, and subsequent designs and analyses were done using our improved CPD programs PHOENIX and TRIAD. Details are provided in SI Methods.

Protein Expression and Purification. All proteins were expressed using BL21 DE3 cells transformed by pET plasmids with 1 mM isopropyl β-D-1-thiogalactopyranoside in standard LB at 16 °C (His6-ENH and ENH-c2a-Strep) or 37 °C (all other proteins). The 13C/15N-labeled ENH-c2b-Strep for NMR experiments was prepared by growing BL21 DE3 cells in 1 L of LB until the OD600 reached ~0.6 and transferring the cells to 250 mL of M9 medium with 13C-glucose and 15N ammonium chloride. Purification of ENH-c2b was accomplished by fusing it to His6-ubiquitin, running the construct on an Ni2+ -NTA column (Qiagen), and then cleaving His6-ubiquitin off using ubiquitin carboxyl-terminal hydrolase isozyme L3 protease (37 °C overnight). Strep-Tactin Sepharose (IBA) and Superdex 75 (Amersham Pharmacia) columns were used for Strep-tag affinity chromatography and size exclusion chromatography, respectively. Details are provided in SI Methods.

CD Spectroscopy. CD studies were performed on an Aviv 62A DS spectropolarimeter equipped with a thermostatic temperature controller. Samples were prepared in 100 mM sodium chloride and 20 mM sodium phosphate buffer at pH 7.5. Wavelength scans and temperature denaturations were carried out in cuvettes with a 0.1-cm pathlength at a protein concentration of ~10 μM. Three wavelength scans were performed at 25 °C for each sample and averaged. The thermal denaturation curve was collected at 222 nm from 0 to 99 °C, sampling every 1 °C separated by 2-min equilibration times (signal averaging time was 1 s). The refolding curve was collected after the thermal denaturation experiment using the same sample.

Analytical Ultracentrifugation. ENH-c2b was analyzed on an XL-1 analytical ultracentrifuge equipped with an AnTi60 rotor (Beckman Coulter). Two-channel Epon-filled centerpieces were used for the sedimentation velocity experiment. Cells were torqued to 130 pounds per inch and run at 60,000 rpm. Data were acquired at 230 nm and 20 °C in continuous mode. Data were first fit to the c(s) model (continuous distribution of sedimentation coefficient) and then converted to the c(M) model (continuous distribution of molecular mass). Time-invariant noises and baseline offsets were corrected before fitting. A maximum entropy regularization confidence level of 0.95 was used in all of the size distribution analyses.

Fluorescence Polarization Assay. Fluorescence polarization was measured at room temperature with a Fluorolog-3 spectrofluorometer (HORIBA). ENH-c2b was serially diluted in buffer containing 100 mM NaCl and 20 mM Tris-HCl at pH 8.0. Fluorescence anisotropy was measured for each sample, and the G-factor was determined individually. Data were analyzed according to a simple monomer-dimer equilibrium model and fit with KaleidaGraph software (Synergy Software). Polarization values for the completely monomeric and dimeric states were fit to be 251 mA and 12 mA, respectively.

![Fig. 5](image-url) Comparison of ENH-c2b averaged minimized solution NMR structure (green) and design model (gray). (A) Superposition of a single chain from the NMR structure and design model. (B) Alternative view of A: ~180° of rotation about the vertical axis. (C) Superposition of the entire dimer NMR structure and design model. (D) Alternative view of C: ~90° of rotation about the horizontal axis. (E) Superposition of the left chain of the NMR structure with the left chain of the design model showing the entire dimer structure. (F) Alternative view of E: ~90° of rotation about the horizontal axis. N, N terminus.

![Fig. 6](image-url) NMR spectrum showing intermolecular NOE restraints obtained by a 3D 13C/15N-filtered NOESY-H-15C-HOSQC experiment. Contour plots of [α1(NH)] and [α1(NH)]-strips of Ala20C, Ala16C, Leu39C, and Leu199 are shown. Chemical shifts indicated on the top and bottom correspond to α1(NH) and α1(NH) dimensions, respectively. For clarity, only the aliphatic region in the α1(NH) dimension is shown. Unambiguous restraints identified for Ala16, Ala20, Leu19, and Leu39 residues are labeled.
**Solution NMR Experiments.** All spectra were acquired at 310 K on a Bruker Avance III 800 spectrometer equipped with a 5-mm z-gradient TCI (H, 13C, and 15N) CryoProbe (Bruker). ENH-2b-Strap (1.9 mM protein in 300 μL) was dissolved in 100 mM NaCl, 5 mM CaCl2, 10 mM DTT, 0.02% NaN3, 5% D2O, and 20 mM NH4OAc at pH 4.5 in a Shigemi NMR tube. Assignment of main-chain and side-chain chemical shifts was based on 1H-15N HSQC, 1H-13C HSQC, CBCA(CO)NH, HNCA, HNCC, HCC-COSY, HCH-TOCSY, HBHA(NH), HNHA(CO)NH, (H)CC(CO)NH, HNHA, CACO, CON, and 13N-TOSY-HSQC experiments. NOE distance restraints were obtained from 1H-15N-edited NOESY, 1H-13C-edited NOESY (aliphatic), and 15C-edited NOESY (aromatic) for intrachain or interchain contacts. An asymmetrically labeled dermer was prepared by mixing 1:1 uniformly 13C/15N-labeled and unlabeled ENH-2b. This sample was used for the 3D 13C/15N-filtered NOESY-1H-13C-HSQC experiment (Fig. S4) to extract the interchain NOE restraints (S3, S4). Details, including structure determination, are described in SI Methods.

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