Design and characterization of a protein fold switching network

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Abstract

Protein sequences encoding three common small folds $(3\alpha, \beta\text{-grasp}, \text{and } \alpha/\beta\text{-plait})$ were connected in a network with high-identity intersections, termed nodes. The structures of proteins around nodes were determined using NMR spectroscopy and analyzed for stability and binding function. To generate nodes, the amino acid sequence encoding a shorter fold $(3\alpha \text{ or } \beta\text{-grasp})$ is embedded in the structure of the ~50% longer α/β -plait fold and a new sequence is designed that satisfies two sets of native interactions. This leads to protein pairs with a 3α or β -grasp fold in the shorter form but an α/β -plait fold in the longer form. Further, embedding smaller antagonistic folds in longer folds creates critical states in the longer folds such that single amino acid substitutions can switch both their fold and function. This suggests that abrupt fold switching may be a mechanism of evolving new protein structures and functions.

INTRODUCTION

Fold switching occurs when one amino acid sequence has a propensity for two completely different, but well-ordered, conformations. Many examples of both natural and engineered fold switching demonstrate that proteins can have a stable native fold while simultaneously hiding latent propensities for alternative states with new functions ¹⁻⁸. This fact has many implications for understanding how amino acid sequence encodes structure, how proteins evolve, how mutation is related to disease, and how function is annotated to sequences of unknown structure ⁹⁻²².

In this work, a network of high-identity intersections (nodes) was engineered that connects three common and well-studied protein folds. Two of these folds are from Streptococcal Protein G which contains two types of domains that bind to serum proteins in blood: the G_A domain binds to human serum albumin (HSA) 23,24 and the G_B domain binds to the constant (Fc) region of IgG 25,26 . The third protein is S6, a component of the 30S ribosomal subunit of *Thermus thermophilus* $^{27-31}$. For simplicity, the S6 fold is referred to as an S-fold, the G_A fold as an A-fold, and the G_B fold as a B-fold. These proteins share no significant homology and are representative of three of the ten most common folds: the S-fold is a thioredoxin-like α/β plait; the A-fold is a homeodomain-like 3α —helix bundle; the B-fold is a ubiquitin-like β grasp³².

Fig. 1 depicts a map of the nodes that were engineered here. The arrows in Fig. 1 show a network originating with the natural S6 sequence. Circles represent nodes in the network at which structural and/or functional switches occur. The SI and S'I nodes are a branch point and lead down diverging sequence pathways, one leading to a node with the A-fold (S/A) and one to a node with the B-fold (S/B). Intersecting mutational pathways lead from S/A to the native G_A protein and S/B to the native G_B protein. At this intersection (A/B), an A-fold switches to a B-fold.

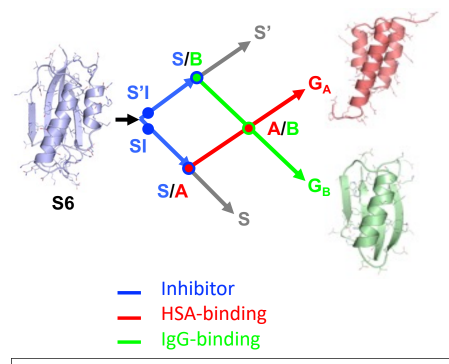


Fig. 1: Overview of engineered nodes in the S6, G_A , and G_B network. S6 is the origin sequence in the engineering process. SI and S'I are separate nodes and are loop size variants of the S-fold, both having protease inhibitor function. The SI branch of the mutational path leads to a node with the A-fold and HSA binding function. The S'I branch of the path leads to a node with the B-fold and IgG binding function. The S/A node (blue and red circle) includes proteins S_{a1} , S_{a2} , A_1 , and A_2 . The S/B node (blue and green circle) includes proteins S_{b3} , S_{b4} , S_{b5} , S_{a3} , and S_{b4} . The A and B paths themselves intersect at an A/B node (green and red circle) at which A- and B-folds are nearly isoenergetic and bifunctional. The S and S' branches continue and connect with many other natural sequences in the α/β plait super-fold family.

Proteins around the A/B node have been extensively characterized in our earlier work and demonstrated protein structure can be encoded by a small number of essential residues, and that a very limited subset of intra-protein interactions can tip the balance from one fold and function to another $^{33-35}$. Here we determine that both G_A and G_B can switch into a third fold $(\alpha/\beta-\text{plait})$ and show that three folds and four functions can be connected in a network that avoids unfolded and functionless states. We describe how these nodes were engineered, determine key structures using NMR spectroscopy, and analyze stability and binding function. The ability to design and characterize nodes connecting three common small folds suggests that fold switching may be a general phenomenon in the evolution of protein structure and function.

RESULTS

Designing a functional switch from ribosomal protein to protease inhibitor. The S6 ribosomal protein is structurally homologous to subtilisin protease inhibitors known as prodomains (Fig. 2A, B) 36 . Prodomain-type inhibitors have two binding surfaces with the protease. One surface comprises the last nine C-terminal amino acids of the inhibitor which bind in the substrate binding cleft of the protease (Fig. 2B). A second, more dynamic surface is formed between two subtilisin helices and the large surface of the β–sheet in the α/β -plait topology of the inhibitor (Fig. 2B) $^{37-39}$. As a result, the S6 protein could be converted into a subtilisin inhibitor protein of the same overall fold (denoted SI) by replacing its nine C-terminal amino acids with residues optimized to bind in the substrate binding cleft of subtilisin. This replacement results in new contacts between the SI β–sheet and the subtilisin surface helices (Fig. 2B).

The SI-protein is 99 amino acids in length and has a 10 residue loop between $\beta 2$ and $\beta 3$. However, there are many natural variations in the length of loops in the conserved α/β -

plait topology ⁴⁰. Therefore, we also engineered a 91 amino acid version of the S-fold (denoted S'I), which resembles the topology of natural prodomain inhibitors (**Fig. S1**). Specifically, the S'I inhibitor has a longer loop connecting $\beta 1$ to $\alpha 1$ and a shorter turn connecting $\beta 2$ to $\beta 3$ (**Fig. 2B**).

The SI and S'I proteins were expressed and purified by binding to a protease column ⁴¹. The CD spectra were compared to the native S6 protein (**Fig. S1**). Inhibition constants (K_I) were measured using an engineered RAS-specific subtilisin protease and the peptide substrate QEEYSAM-AMC ⁴¹. SI and S'I inhibit the RAS-specific protease with K_I values of 200nM and 60nM, respectively (**Table S1**). The details of the competitive inhibition assay are described in Materials and Methods. The results demonstrate that a ribosomal protein can be converted into a protease inhibitor with minor modification (and without a fold switch). In addition, however, the SI and S'I proteins also facilitated engineering subsequent switches to new folds and functions by linking each of the S-, A-, and B- folds to easily measured binding functions: protease inhibition (S or S'-fold); HSA-binding (A-fold); and IgG binding (B-fold).

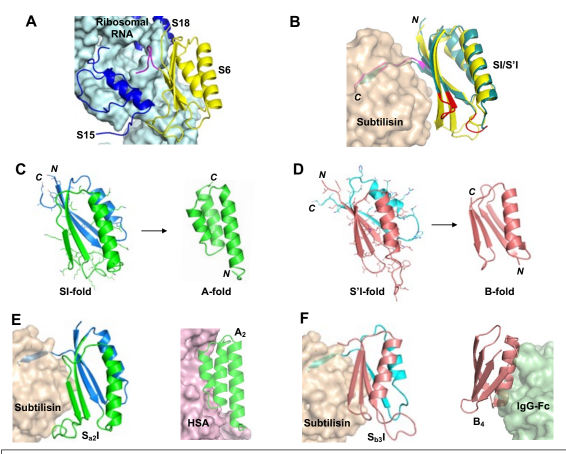


Fig. 2: Summary of switches in structure and function. A) Structure of the S6 protein (yellow), RNA (light blue), and S15 and S18 proteins (blue) in the 30S ribosome (PDB 1FKA). C-terminal amino acids of S6 are in magenta. B) Subtilisin (wheat) is shown in complex with a model of the SI-inhibitor (yellow). The C-terminal nine amino acids of SI are shown in magenta. These positions were mutated in native S6 to generate affinity for subtilisin. The S'I-inhibitor (teal) is also shown with the altered loops in red. The subtilisin used in the modeling was the engineered RAS-specific protease. C) The Sa1 protein (blue and green) was generated from SI by mutating the 45 positions (mutant side chains shown with sticks). Deletion of C-terminal amino acids (blue) switches Sa1 into the A-fold (green). D) The Sb3 protein (rose and cyan) was generated from S'I by mutating the 67 positions (mutant side chains shown with sticks). Deletion of C-terminal amino acids (cyan) or point mutation will switch Sb3 into a B-fold (rose). E) Model of Sa2 (green and blue) bound to subtilisin (wheat). Model of A2 (based on A1 structure) bound to HSA (violet). The HSA complex used PDB 2VDB as the template. F) Model of Sb3 (rose and cyan) bound to subtilisin (wheat). Model of B4 (rose) bound to Fc (mint). The Fc complex used PDB 1FCC as the template. The subtilisin used in the modeling and inhibition measurements was the engineered RAS-specific protease (PDB 6UAO).

Designing fold switches. In previous work we created sequences that populate both A- and B-folds by threading the A-sequence through the B-fold, finding a promising alignment, and then using phage-display selection to reconcile one sequence to both folds ^{34,42,43}. Here the

approach is conceptually similar, except that we use Rosetta ⁴⁴ as a computational design tool to test compatible mutations rather than phage display. The design process is as follows:

- i. Thread the A- or B- sequence through both SI and S'I-fold types.
- ii. Identify alignments that minimize the number of catastrophic interactions.
- iii. Design mutations to resolve unfavorable interactions in clusters of 4-6 amino acids using Pymol ⁴⁵ and energy minimize using Rosetta-Relax ⁴⁴.
- iv. Optimize protein stability in the S-fold by computationally mutating amino acids at nonoverlapping positions. Repeat energy minimization and evaluation with Rosetta-Relax.
- v. To reduce uncertainties involved in computational design, conserve original amino acids whenever possible.

There is no reason to assume that this method is optimal. We are just applying a practicable scheme for engineering sequences compatible with two sets of native interactions and then evaluating structure, stability, and function. Initial designs were refined based on structural analysis with NMR, thermodynamic analysis of unfolding, and functional analysis using binding assays, as described below. All designed proteins were expressed in *E. coli* and purified to homogeneity as described in Materials and Methods.

Designing a switch from α/β -plait protease inhibitor to 3α HSA-binding protein. Alignment of the 56 amino acid HSA-binding, A-fold with the 99 amino acid SI-fold and subsequent mutation to resolve catastrophic interactions produced low energy switch candidates denoted S_{a1} and A_1 . The exact sequence of A_1 is embedded in S_{a1} at positions 11-66 such that the $\alpha 1$ helices are structurally aligned (Fig. 3A, Fig. S2A). Their final computational models were generated by Rosetta using the Relax application. The Relax protocol searches the local conformational space around an experimentally-determined structure and is used only to evaluate whether the designed mutations have favorable native interactions within that limited

conformational space. The designed models of S_{a1} and A₁ are very similar in energy compared to the respective relaxed native structures (**Fig. S3 and Supplemental PDB files of the Rosetta models**).

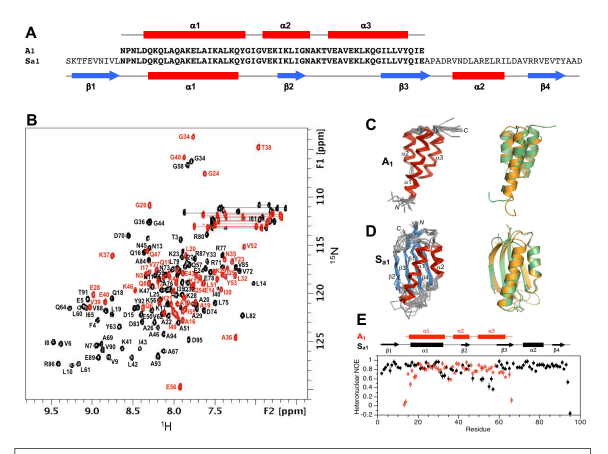


Fig. 3: Structure and dynamics of A_1 and S_{a1} . (**A**) Sequence alignment of A_1 and S_{a1} , which are 100% identical over the 56 amino acid A-region. (**B**) Overlaid two dimensional ${}^{1}H^{-15}N$ HSQC spectra of S_{a1} (black) and A_1 (red) with backbone amide assignments. Spectra were recorded at 25°C and 5°C, respectively. (**C**) Ensemble of 10 lowest energy CS-Rosetta structures for A_1 (left panel). Superposition of the A_1 structure (green) with the parent G_A fold (orange) (right panel). (**D**) Ensemble of 10 lowest energy CS-Rosetta structures for S_{a1} (left panel). Superposition of S_{a1} (green) with the parent S6 fold (orange) (right panel). (**E**) Backbone dynamics in designed proteins. Plot of ${}^{1}H_{1}^{-15}N$ steady state heteronuclear NOE values at 600 MHz versus residue for A_1 (red) and for S_{a1} (black). Error bars indicate ±1SD.

Structural analysis of A_1 and S_{a1} . Overall, the 3α -helical bundle topology of A_1 is very similar to the G_A parent structure from which it was derived ⁴⁶. The sequence specific chemical shift assignments for A_1 (**Fig. 3B**) were utilized to calculate a 3D structure with CS-Rosetta (**Fig. 3C**, **Table 1**). Our previous studies indicated close correspondence of CS-Rosetta and *de novo* structures for A- and B-folds with high sequence identity ⁴⁷. The N-terminal residues 1-4 and the

C-terminal residues 53-56 are disordered in the structure, consistent with ${}^{1}H{}^{-15}N$ steady state heteronuclear NOE data (**Fig. 3E**). Likewise, S_{a1} has the same overall $\beta\alpha\beta\beta\alpha\beta$ -topology as the parent S6 structure (**Fig. 3D**, **Table 2**). The backbone chemical shifts (**Fig. 3B**) were used in combination with main chain inter-proton NOEs (**Fig. S4**) to determine a three-dimensional structure utilizing CS-Rosetta (**PDB 7MN1**). The conformational ensemble shows well-defined elements of secondary structure at residues 2-10 (β 1), 16-32 (α 1), 40-44 (β 2), 59-67 (β 3), 73-81 (α 2) and 86-92 (β 4). The principal difference from the native structure is that the β 2-strand is seven amino acids shorter in S_{a1} than in S6. Heteronuclear NOE data show overall consistency with the structure, indicating that the long loop between the β 2- and β 3-strands from residues 45-58 is more flexible than other internal regions of the polypeptide chain (**Fig. 3E**).

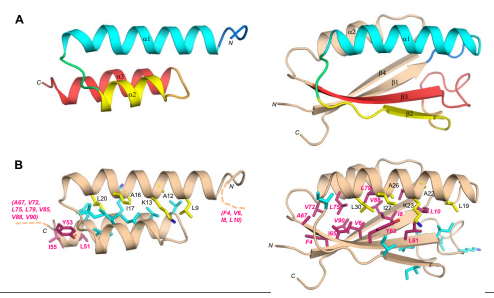


Fig. 4: Structural differences between the 100% sequence identical regions of A_1 and S_{a1} . (A) Main chain comparisons. (Left panel) CS-Rosetta structure of A_1 with color coding for secondary structured elements. (Right panel) Corresponding color-coded regions mapped onto the CS-Rosetta structure of S_{a1} , illustrating changes in backbone conformation. Regions outside the 56 amino acid sequence of A_1 are shown in wheat. (B) Side chain comparisons. (Left panel) Residues contributing to the core of A_1 from the $\alpha 1$ -helix (yellow), and from other regions (cyan). The non- $\alpha 1$ core residues from S_{a1} (pink) do not overlap with the A_1 core (see text for further details). (Right panel) Residues contributing to the core of S_{a1} from the $\alpha 1$ -helix (yellow), and most of the other participating core residues (pink). The non- $\alpha 1$ core residues from A_1 are also shown (cyan), highlighting the low degree of overlap.

Comparison of A_1 and S_{a1} structures. Although the 56 amino acid sequence of A_1 is 100% identical to residues 11-66 of Sa1, a significant fraction of the backbone undergoes changes between the two structures. Most notably, while the $\alpha 1$ helices in both A_1 and S_{a1} are similar in length, the regions corresponding to the α 2 and α 3 helices of A₁ form the β 2 and β 3 strands of S_{a1} (Fig. 4A). Core amino acids in the α 1-helix of A_1 correspond with residues that also contribute to the core of S_{a1} . However, the $\alpha 1$ -helix in S_{a1} contacts an almost entirely different set of residues (Fig. 4B). For example, amino acids L51, Y53, and I55 in the C-terminal tail of A₁ do not have extensive contacts with $\alpha 1$ but the corresponding residues in S_{a1} (L61, Y63, and I65) form close core interactions with $\alpha 1$ as part of the $\beta 3$ -strand. Most of the other core residues contacting the α 1-helix of S_{a1} are outside the 56 amino acid region coding for the A₁ fold. These include F4, V6, I8, and L10 from the β1-strand; A67 from the β3-strand; V72, L75, and L79 from the α 2-helix; and V85 from the loop between the α 2-helix and the β 4-strand. Two additional residues, V88 and V90 (β 4) also contribute significantly to the core but do not contact α 1. Thus, except for the original topological alignment of the α 1-helices, the cores of the 3α and α/β -plait folds are largely non-overlapping. In total, approximately half of the residues participating in the S_{a1} core are not present in the A_1 sequence.

Energetics of unfolding for A_1/S_{a1} . Far-UV CD spectra were measured for S_{a1} and A_1 and their thermal unfolding profiles were determined by measuring ellipticity at 222nm versus

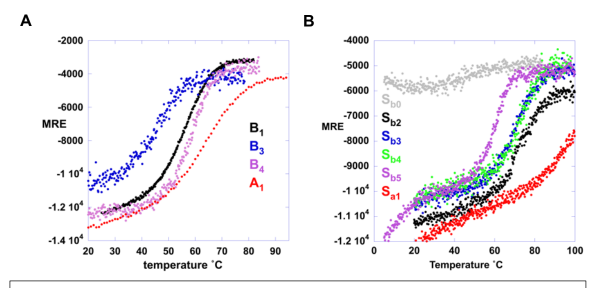


Fig. 5: CD melting curves. (A) Ellipticity at 222nm plot versus temperature for A- and B- variations. (B) Ellipticity at 222nm plot versus temperature for S-variations. S_{b0} is a low-stability variant (F7V) of S_{b1} used to measure the temperature dependence of the unfolded state.

temperature (**Fig. 5 and S5**). S_{a1} has a T_M of ~100°C and an estimated $\Delta G_{folding}$ of -5.3 kcal/mol at 25°C (**Fig. 5B**, **Table S1**) ⁴⁸. The $\Delta G_{folding}$ of the parent S6 is -8.5 kcal/mol ³⁰. The Rosetta energy of the S_{a1} design is similar to that of the native sequence (**Fig. S3**). A_1 has a T_M of 65°C and a $\Delta G_{folding} = -4.0$ kcal/mol at 25°C ⁴⁸ (**Fig. 5A**, **Table S1**). The $\Delta G_{folding}$ of the parent G_A is -5.6 kcal/mol ^{49,50}. The Rosetta energy of the A_1 design is slightly more favorable than for the native sequence (**Fig. S3**).

HSA binding. Initial engineering of the fold switch was carried out without consideration of preserving function. As a result, A_1 does not have detectible HSA binding affinity because two amino acids in the binding interface were mutated. Significant HSA-binding is recovered, however, when the surface mutations, E28Y and K29Y, are made in A_1 (denoted A_2). These mutations do not appear to affect the structure of A_1 (**Fig. S5**) but result in HSA binding of $K_D \le$

 $1\mu M$ (**Table S1**). This was determined by measuring binding to immobilized HSA as described in Material and Methods.

Protease inhibition. S_{a1} does not bind protease because C-terminal amino acids were not preserved in its design. It can be converted into a protease inhibitor, however, by replacing its three C-terminal amino acids (AAD) with DKLYRAL (denoted $S_{a1}I$). A version of $S_{a1}I$ was also made that contains the exact 56 amino acid A_2 sequence by making E38Y, K39Y mutations (denoted $S_{a2}I$). S_{a1} , $S_{a1}I$, and $S_{a2}I$ are similar in structure by CD analysis (**Fig. S5**). The inhibition constant of $S_{a2}I$ with the engineered subtilisin was determined to be 50nM as described in Materials and Methods (**Table S1**). Thus, a stable A-fold with HSA-binding function can be embedded within a 99 amino acid S-fold with protease inhibitor function (**Fig. 2C and E**). It should be noted that all HSA contact amino acids are preserved in both the A_2 and $S_{a2}I$ sequences, but the three-dimensional topology necessary to form the HSA contact surface occurs only in the A-fold. Nevertheless, $S_{a2}I$ was observed to bind weakly to HSA ($K_0 \sim 100 \mu M$), **Table S1**). This weak affinity suggests that some $S_{a2}I$ molecules may populate the 3α fold even though the α/β -plait fold strongly predominates.

Designing a switch from α/β -plait protease inhibitor to β -grasp IgG-binding protein. In designing an S- to B-fold switch, we used two topological alignments. The first was between SI- and B-folds, where the $\beta 1$ strands of each fold were aligned (Fig. S2B, Fig. S6A). The second alignment was between S'I- and B-folds, where the long loop between $\beta 2$ and $\beta 3$ in SI was shortened in S'I to be more consistent with natural protease inhibitors. In this scheme, the $\alpha 1\beta 3\beta 4$ topology of the B-fold was aligned with the $\alpha 1\beta 2\beta 3$ topology of the S'I-fold (Fig. 6A, Fig. S2C).

Design and characterization of B_1 , S_{b1} , B_2 , and S_{b2} . In the first approach, alignment of the β 1-strands of the B-fold and the S-fold and subsequent mutation to resolve catastrophic

interactions produced low energy switch candidates denoted B_1 and S_{b1} . The exact sequence of B_1 is embedded in S_{b1} at positions 4-59 (Fig. S6A). The computational models of B_1 and S_{b1} show relatively small increases in energy compared to the corresponding relaxed native structures (Fig. S3). The NMR structure of B1 displayed a $\beta\beta\alpha\beta\beta$ topology identical to that of the parent B-fold, with a backbone RMSD of ~0.6 Å (Fig. S6B, C). The topology of S_{b1} is not the same as the parent S6 structure, however, and instead has a fold similar to that of B_1 (Fig. S6B ,D; Fig. S7, PDB 7MQ4). Introducing 13 mutations into S_{b1} generated a protein denoted S_{b2} (Fig. S8). S_{b2} contains four β -strands and two α -helices and has the general features of the parent S-fold (Fig. S9, PDB 7MN2). The 56 amino acid version of S_{b2} (denoted B_2) has a significantly higher Rosetta energy than B_1 , however, and is presumably unfolded (Fig. S3). Thus, neither the B_1/S_{b1} nor B_2/S_{b2} protein pairs resulted in high identity sequences with different folds. Nonetheless, B_1 is 80% identical to the corresponding embedded region in the S-folded protein S_{b2} (Fig. S9A). The structures of B_1 , S_{b1} , and S_{b2} are described further in the Supplement and Tables 1 and 2.

Design of S_{b3} **and** B_3 . To improve the design of the S to B switch we aligned the B-fold with the S' inhibitor fold and chose an alignment that creates a topological match between $\alpha 1\beta 3\beta 4$ in B and $\alpha 1\beta 2\beta 3$ in S' (**Fig. S2C**). Mutation to resolve deleterious interactions in this alignment produced low energy switch candidates denoted B_3 and S_{b3} (**Fig. S10**). The exact sequence of B_3 is embedded in S_{b3} at positions 1-56. The energy of the computational model for S_{b3} is slightly more favorable than the relaxed native structure. The designed model of B_3 shows relatively small increases in energy compared to the relaxed native structure (**Fig. S3**).

Structural analysis of S_{b3} and B_3 . NMR-based structure determination indicated that S_{b3} has a $\beta\alpha\beta\beta\alpha\beta$ secondary structure and an S-fold topology (Fig. 6A,B,D, PDB 7MP7). Ordered regions correspond with residues 4-10 (β 1), 24-37 (α 1), 42-46 (β 2), 51-56 (β 3), 62-70 (α 2), and

79-85 (β 4). Comparison of S_{b3} with the parent S-fold indicates that the β 1/ α 2/ β 4 portion of the fold is similar in both. In contrast, the β 1- α 1 loop is longer in S_{b3} (13 residues) than in the parent S-fold (5 residues), while α 1, β 2, the β 2- β 3 loop, and β 3 are all shorter than in the parent (**Fig. 6D**). Consistent with the S_{b3} structure, the 13 amino acid β 1- α 1 loop is highly flexible (**Fig. 6E**). We also expressed and purified a truncated protein corresponding to the embedded B-fold, the 56 amino acid version of S_{b3} (denoted B_3). The 2D 1 H- 1 5N HSQC spectrum of B_3 at 5°C and low concentrations (<20 μ M) was consistent with a predominant, monomeric B-fold (**Fig. S11**) but showed significant exchange broadening at 25°C, indicative of low stability (see below). Presumably the low stability is due to less favorable packing of Y5 in the core of the B-fold compared with a smaller aliphatic leucine. However, additional, putatively oligomeric, species were also present for which relative peak intensities increased with increasing protein concentration. Due to its relatively low stability and sample heterogeneity, B_3 was not analyzed further structurally.

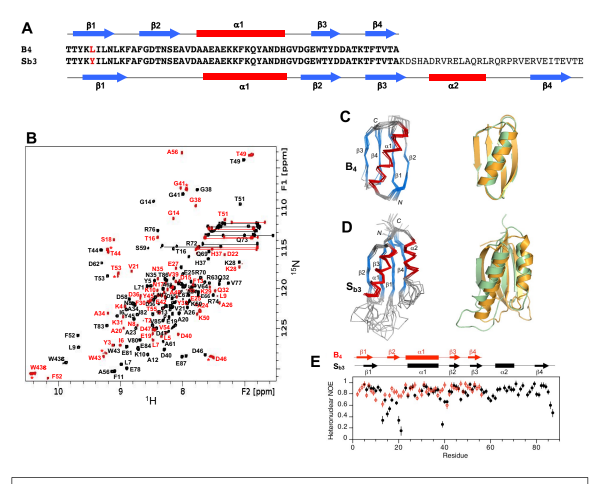


Fig. 6: Structure and dynamics of S_{b3} and B_4 . (A) Sequence alignment of B_4 and S_{b3} , differing by one residue (L5Y) over the 56 amino acid B-region. (B) Overlaid two dimensional ${}^1H^{-15}N$ HSQC spectra of S_{b3} (black) and B_4 (red) with backbone amide assignments. Spectra were recorded at 25°C. The A56 peak is an aliased signal. Peaks labeled with an asterisk decrease in relative intensity as the B_4 concentration is lowered, indicating the presence of a weakly associated putative dimer in addition to monomer. (C) Ensemble of 10 lowest energy CS-Rosetta structures for B_4 (left panel). Superposition of the B_4 structure (green) with the parent G_B fold (orange) (right panel). (D) Ensemble of 10 lowest energy CS-Rosetta structures for S_{b3} (left panel). Superposition of S_{b3} (green) with the parent S6 fold (orange) (right panel). (E) Plot of 1H }- ${}^{15}N$ steady state heteronuclear NOE values at 600 MHz versus residue for B_4 (red) and S_{b3} (black). Error bars indicate ±1SD.

Design and analysis of point mutations that switch the fold of S_{b3} . We used the NMR structure of S_{b3} to design a point mutation, tyrosine 5 to leucine (Y5L), that would stabilize the embedded B-fold without compromising native contacts in the S-fold (**Fig. S10**). This mutant was therefore expected to shift the population to the B-fold. The Y5L mutant of S_{b3} (denoted S_{b4}) exhibited approximately twice the number of amide cross-peaks in the HSQC spectrum relative to the S_{b3} sample. Comparison of the spectrum of S_{b4} with spectra of S- and B-folds for closely

related sequences indicated that S_{b4} populates both S- and B-states simultaneously in an approximately 1:1 ratio at 25°C (**Fig. S12**). A complete NMR analysis of the kinetics of interconversion will be reported elsewhere. The Y5L mutation also was introduced into B_3 to determine its effect on the stability of the B-fold in the 56 amino acid protein. The new protein, denoted B_4 , is indeed more stable than B_3 (**Fig. 5A, Table S1**). Assignment and structure determination of B_4 showed its topology to be identical to the parent B-fold (**Fig. 6B, C**). At concentrations above 100 μ M, B_4 displayed a tendency for weak self-association similar to that seen for B_3 .

Finally, we designed a mutation of leucine 67 to arginine (L67R) in S_{b4} to destabilize the S-fold without changing the sequence of the embedded B-fold. The mutant is denoted as S_{b5} (**Fig. S10**). This was expected to shift the population to the B-fold. The 2D $^1H^{-15}N$ HSQC spectrum of S_{b5} indicates that the L67R mutation does indeed destabilize the S-fold, with the loss of S-type amide cross-peaks and the concurrent appearance of a new set of signals indicating a switch to a B-fold. Superposition of the spectrum of S_{b5} with that of B_4 shows that the new signals in S_{b5} largely correspond with the spectrum of B_4 (**Fig. S13**). Thus, the L67R mutation shifts the equilibrium from the S-fold to the B-fold. The additional signals (~25-30) in the central region of the HSQC spectrum that are not detected in B_4 are presumably due to the disordered C-terminal tail of S_{b5} . The C-terminal tail of S_{b5} does not appear to interact extensively with the B-fold, as evidenced by few changes in chemical shifts or peak intensities in the B-region of S_{b5} compared with B_4 .

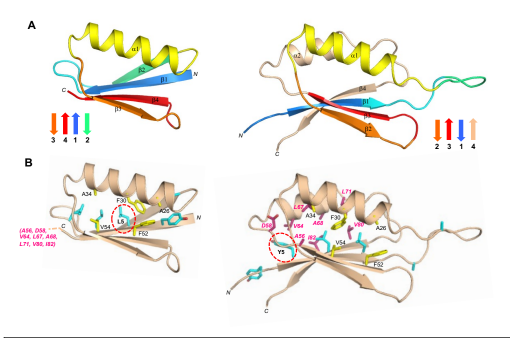


Figure 7: Structural differences in the high (~98%) sequence identity regions of B_4 and S_{b3} . (A) Main chain comparisons. (Left panel) CS-Rosetta structure of B_4 with secondary structure elements color coded. (Right panel) Corresponding color-coded regions mapped onto the CS-Rosetta structure of S_{b3} , showing changes in backbone conformation. Regions outside the 56 amino acid sequence of B_4 are shown in wheat. (B) Side chain comparisons. (Left panel) Residues contributing to the core of B_4 from $\alpha 1/\beta 3/\beta 4$ (yellow), and from other regions (cyan). The non- $\alpha 1/\beta 2/\beta 3$ core residues from S_{b3} (pink) do not overlap with the B_4 core (see text for further details). (Right panel) Residues contributing to the core of S_{b3} from $\alpha 1/\beta 2/\beta 3$ (yellow), and the other participating core residues (pink). The non- $\alpha 1/\beta 2/\beta 3$ core residues from B_4 are also shown (cyan).. The single L5Y amino acid difference between B_4 and S_{b3} is highlighted.

Structural comparison of S_{b3} and B_4 . The aligned amino acids 1-56 of S_{b3} and B_4 have 98% sequence identity, the only difference being an L5Y mutation in S_{b3} (Fig. 6A). The global folds of S_{b3} and B_4 have large-scale differences, however (Fig. 7A, Fig. S4). The β 1-strands, while similar in length, are in opposite directions in S_{b3} and B_4 . The β 1-strand forms a parallel stranded interaction with β 4 in B_4 , but an antiparallel interaction with the corresponding β 3-strand in S_{b3} . Whereas residues 9-20 form the 6-residue β 1- β 2 turn and the 6-residue β 2-strand of B_4 , these same amino acids constitute the end of β 1 and 10 residues of the large disordered β 1- α 1 loop in S_{b3} . The remainder of the B-region is topologically similar, with the α 1/ β 3/ β 4 structure in B_4 matching the α 1/ β 2/ β 3 structure in S_{b3} . Overall, however, the order of H-bonding in the 4-stranded β -sheets is quite different, with β 2 β 3 β 1 β 4 in S_{b3} and β 3 β 4 β 1 β 2 in B_4 .

The main core residues of B_4 consist of Y3, L5, L7, and L9 from $\beta 1$, A26, F30, and A34 from $\alpha 1$, and F52 and V54 from $\beta 4$ (**Fig. 7B**). In S_{b3} , the topologically equivalent regions of the core are A26, F30, and A34 from $\alpha 1$, and F52 and V54 from $\beta 3$. Residues Y5, L7, and L9 from the $\beta 1$ strand of S_{b3} also form part of the core, but with different packing from B_4 due to the reverse orientation of $\beta 1$. Residues A12 and A20, which contribute to the periphery of the core in B_4 , are solvent accessible in the $\beta 1$ - $\alpha 1$ loop of S_{b3} . Most of the remaining core residues of S_{b3} come from outside of the B-region and include amino acids from $\beta 3$ (A56), $\alpha 2$ (V64, L67, A68, L71), and $\beta 4$ (V80 and I82).

Energetics of unfolding for B_3/S_{b3} , B_4/S_{b4} , and S_{b5} . Far-UV CD spectra were measured for B_3 , B_4 , S_{b3} , S_{b4} , and S_{b5} and their thermal unfolding profiles were determined by measuring ellipticity at 222nm versus temperature (Fig. 5, S10, Table S1). As described above, the predominant form of S_{b3} is an S-fold. CD and NMR analysis show that B_3 is predominantly a B-fold with a $\Delta G_{folding}$ of -1.2 kcal/mol at 25°C ⁴⁸. From the NMR analysis, it appears that the B-fold is in equilibrium with putatively dimeric states. This creates a situation in which the B-fold is both temperature dependent and concentration dependent. The predominant form at 5°C and $\leq 18\mu M$ is the B-fold, however. The low stability and concentration-dependent behavior of B_3 may indicate that some propensity for S-type conformations could persist in the 56-residue protein.

 S_{b4} has a temperature unfolding profile very similar to S_{b3} even though both S- and B- are equally populated at 25°C in S_{b4} (**Fig. 5**). This shows that the Y5L mutation results in two folds that are almost isoenergetic and both thermodynamically stable relative to the unfolded state. Further, because S- and B-folds are in equilibrium and equally populated, the free energy of switching to the B-fold from the S-fold ($\Delta G_{B-fold/S-fold}$) is ~ 0 kcal/mol at 25°C. The switch

equilibrium reflects the influence of the antagonistic B-fold on the S-fold population in S_{b4} , where the leucine at residue 5 helps stabilize the alternative B-state at the expense of the S-state. Thermal denaturation by CD shows that B_4 has a $\Delta G_{folding} = -4.1$ kcal/mol at 25° C 48 . The thermal unfolding profile of S_{b5} shows a low temperature transition with a midpoint ~10°C and a major transition with a midpoint of ~60°C (**Fig. 5B**). The NMR analysis indicates that the major transition is unfolding of the B-fold. Thus, the arginine at 67 in S_{b5} makes the B-fold more favorable by making the S-fold unfavorable, consistent with the change in population from mixed to B-fold observed by NMR.

Protease inhibition. The S_{b3} protein is closely related to S'I but lacks inhibitor function because C-terminal amino acids were changed in the design of the switch. It can be converted into a protease inhibitor, however, by altering C-terminal amino acids VTE to DKLYRAL. This mutant is denoted $S_{b3}I$. S_{b3} and $S_{b3}I$ appear similar in structure by CD analysis (**Fig. S10**). The K_I for $S_{b3}I$ with subtilisin was determined to be 50nM (**Table S1**).

IgG binding. Binding to IgG was determined for B_3 and $S_{b3}I$ (Table S1). B_3 and $S_{b3}I$ bound to IgG Sepharose with $K_D \le 1\mu M$ and $10\mu M$, respectively. Presumably $S_{b3}I$ has significant IgG-binding activity because the $\alpha 1\beta 3$ IgG binding surface of the B-fold is largely preserved in the S-fold. Thus, $S_{b3}I$ is a dual-function protein with both IgG-binding and protease inhibitor functions (Fig. 2F).

DISCUSSION

The entire network of intersecting pathways between the S-, A-, and B-folds is summarized in **Fig. 8**. The first node on the pathway is a functional switch from RNA binding protein to protease inhibitor without a fold switch. The α/β plait is a common fold, and proteins with this basic topology include many different functions ³². Engineering the SI and S'I nodes illustrates how protease inhibitor function can arise in the α/β plait topology with a few

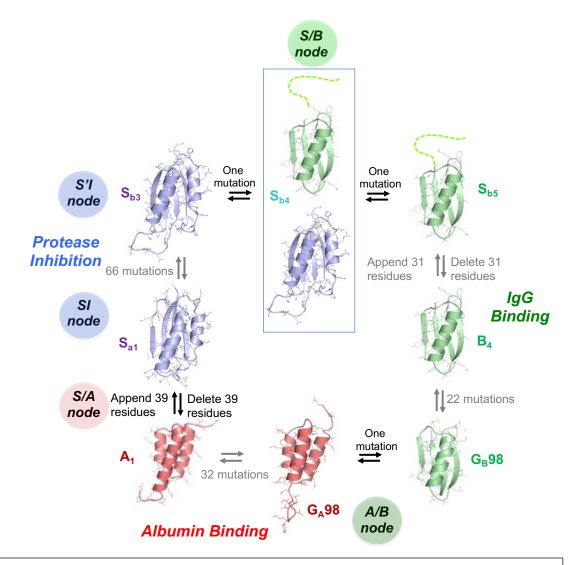


Figure 8: Sequence-fold relationships of engineered S/A, S/B, and A/B nodes. Switches between stable folds can be induced by single amino acid mutation or deleting/appending terminal sequence that stabilizes the S-fold. Blue denotes an S-fold, green a B-fold, and red an A-fold. Gray arrows connect proteins that have been reengineered without a fold switch. S_{b4} is observed with two folds simultaneously. The G_A98 and G_B98 structures are from PDB codes 2LHC and 2LHD, respectively.

mutations. Replacing only C-terminal amino acids in the S6 protein creates interaction with the substrate binding cleft of the protease (**Fig. 2A**, **B**). This C-terminal interaction plus adventitious contact between the β -sheet surface of the α/β plait and two α -helices in the protease results in protease inhibition in the 50nM range. Based on the structure of S6 in the 30S complex, the C-terminal modification may not have major effects on binding interactions with ribosomal RNA and the S15 protein (**Fig. 2A**). Thus, the transition from RNA binding protein to protease inhibitor likely is uninterrupted. An insertion in the β 1- α 1 loop and a deletion β 2- β 3 loop in the SI-inhibitor creates a topology that more closely resembles natural prodomain-type inhibitors^{36,38,51} and creates an α 1 β 2 β 3 motif in the S'-fold that is similar to the α 1 β 3 β 4 motif of the B-fold. This topological similarity brings the S'1 closer to an intersection with the B-fold. Thus, S1 and S'1 nodes are both a functional switch and a branch point for switching the S-fold into the A- and B-folds, respectively.

Engineering nodes at fold intersections required designing sequences that are compatible with native interactions in two different folds. We used simple rules to do this. The first rule was to align topologies rather than maximizing sequence similarities. Identifying a common topology can help determine a register that has fewer irreconcilable clashes. For example, topological alignment of the $\alpha 1$ helix of the SI fold and the $\alpha 1$ helix of the A-fold facilitated engineering the fold switch because the regions flanking $\alpha 1$ of the SI-fold proved to be capable of encoding two different fold motifs. When topological alignment is poor, as was the case with S- and B-folds, it was helpful to look for natural variations in the turns of the longer fold to create better alignment. Variation in loops and turns in a larger fold creates more freedom of design and higher probability of switches. Once an alignment is chosen, the basic rule in resolving catastrophic clashes is to conserve original amino acids when possible. This reduces the uncertainties involved in computational design. The Rosetta energy function was

not used to predict a favorable alignment but was important in evaluating mutations to resolve clashes once an alignment was chosen.

Selecting mutations compatible with two sets of native interactions required tradeoffs in the native state energetics of each individual fold to gain sequence identity 2,8 . A node may be produced in cases in which both alternative folds are stable relative to the unfolded state. Stability relative to the unfolded state (i.e. a state with little secondary structure) was determined by CD melting (**Fig. 5**). It was informative to examine the stability of both short (56 residue) and longer forms of a putative node sequence. The independent stability of the G-fold can be determined in the short form without the antagonism from the S-fold that is present in the longer sequence. The stabilities of the A_1 and A_2 proteins are about -4 kcal/mol at 25° C 48 compared to -5.6 kcal/mol for the native G_A protein 46 . The stabilities of B_3 and B_4 are -1.2 and -4.1 kcal/mol, respectively, at 25° C 48 compared to -6.7 kcal/mol for the native G_B protein 52 . For the longer sequences, the $\Delta G_{\text{folding}}$ of S_{a1} and S_{b3} are -5.3 kcal/mol and -3.5 kcal/mol, respectively, at 25° C 48 compared to -8.5 kcal/mol for the native S6 protein 30 .

In the case of the S-folds, however, the energetic effects of the stable, embedded G-fold must also be considered. Since the equilibria between both folded states and the unfolded state are thermodynamically linked, the free energy of a switch to a G-fold from an S-fold ($\Delta G_{G-fold/S-fold}$) is approximated by the difference in $\Delta G_{folding}$ ($\Delta \Delta G_{folding}$) between the short and long forms of a node protein. For example, based on $\Delta G_{folding}$ for A_1 and S_{a1} , the predicted $\Delta G_{A-fold/S-fold}$ of S_{a1} is 1.3kcal/mol. This is consistent with the structure of the predominant S-fold determined by NMR but also with the small population of 3α fold suggested by weak HSA-binding. From the thermal denaturation profiles of B_3 and S_{b3} , the predicted $\Delta G_{B-fold/S-fold}$ of S_{b3} is 2.3kcal/mol, a value consistent with the stable S-fold observed in NMR experiments. The S_{b3} sequence is also

approaching a critical point, however. A substitution in S_{b3} that stabilizes the B-fold (Y5L) shifts the equilibrium of S_{b4} to an approximately equal mixture of B- and S-folds. That is, $\Delta G_{B\text{-}fold/S\text{-}fold}$ of S_{b4} is ~ 0 kcal/mol at 25°C. One further substitution that destabilizes the S-fold (L67R) shifts the population of S_{b5} to a stable B-fold ($\Delta G_{B\text{-}fold/S\text{-}fold} \le -5$ kcal/mol) (**Fig. 8**).

The existence of nodes between folds has implications for the evolution of new functions. In the case of the S/A node, all contact amino acids for HSA exist within the S-fold of the protease inhibitor $S_{a2}I$ albeit in a cryptic topology. Deletion of amino acids $67-99~(A_2)$ results in loss of inhibitor function and a fold switch from α/β plait to 3α . Acquisition of HSA binding activity ($K_D < 1\mu M$) results from unmasking the cryptic HSA binding amino acids via the fold switch (Fig. 2E). This level of binding affinity could be biologically relevant since the concentration of HSA in serum is $> 500\mu M^{53}$. In the case of the S/B node, the $\alpha I\beta 3$ motif contains all IgG contact amino acids and $S_{b3}I$ has some affinity for both IgG ($K_D = 10\mu M$) and protease ($K_I = 50nM$). In this case, the Y5L mutation (S_{b4}) or a deletion of $57-91~(B_4)$ causes a fold switch from α/β plait to the β –grasp and results in tighter IgG binding ($K_D \le 1\mu M$) (Fig. 2F). This level of binding affinity could also be biologically relevant since the concentration of IgG in serum is $>50\mu M$ (or $>100\mu M$ Fc binding sites) 54 . We have previously shown that an A-fold with HSA binding function can be switched to a B-fold with IgG-binding function via single amino acid substitutions that switch the folds and unmask cryptic contact amino acids for the two ligands 34,35

In conclusion, it was possible to connect three common folds in a network of highidentity nodes that form critical points between two folds. As in other complex systems, a small change in a protein near a critical point can have a "butterfly effect" on how the folds are populated. This property of the protein folding code means that proteins with multiple folds and functions can exist in highly identical amino acid sequences. This suggests that evolution of new folds and functions sometimes can follow uninterrupted mutational pathways.

MATERIALS AND METHODS

Mutagenesis, protein expression and purification. Mutagenesis was carried out using $Q5^{\circ}$ Site-Directed Mutagenesis Kits (NEB). G_A and G_B variants were cloned into a vector (pA-YRGL) encoding the sequence:

MEAVDANSLAQAKEAAIKELKQYGIGDKYIKLINNAKTVEGVESLKNEILKALPTEGSGEEDKQYRGL-

as an N-terminal fusion domain ⁴⁶. Cell growth was carried out by auto-induction ^{34,55}. Cells were harvested by centrifugation at 3,750 x g for 20 minutes and lysed by sonication on ice in 0.1M KPO₄, pH 7.2. Cellular debris were pelleted by centrifugation at 10,000 x g for 15 minutes. Supernatant was clarified by centrifugation at 45,000 x g for 30 minutes. Proteins were purified using a second generation of the affinity-cleavage tag system employed previously to purify switch proteins ^{34,56}. The second-generation tag (*YRGL-tag*) results in high-level soluble expression of the switch proteins and also enables capture of the fusion protein by binding tightly to an immobilized processing protease via the C-terminal EEDQYRGL sequence. Loading and washing were at 5 mL/min for a 5mL *Im-Prot* column using a running buffer of 20mM KPO₄, pH 6.8. The amount of washing required for high purity depends on stickiness of the target protein and how much of it is bound to the column. We typically wash with 10 column volumes (CV) of wash solution followed by 3 CV 0.5M NaCl and then ~ 10 CV running buffer. This can be repeated as necessary. The 0.5M NaCl shots are repeated until the amount of absorbance release with each high-salt shot becomes small and constant. All the high-salt solution is washed out before initiating the cleavage. The target protein was cleaved from the *Im-Prot*

column by injecting 15 mL of imidazole solution (0.1 mM) at 1mL/min, 22°C. The cleaved protein typically elutes as a sharp peak in 2-3 CV. The purified protein was then concentrated to 0.2 to 0.3 mM, as required for NMR analysis. The columns were regenerated by injecting 15 mL of 0.1 N H_3PO_4 (0.227 mL concentrated phosphoric acid (85%) per 100ml) at a flow rate of \sim 1 CV/min. The wash solution was neutralized immediately after stripping. The purification system is available from Potomac Affinity Proteins.

Protease inhibitor proteins were purified by binding to *Im-Prot* media and then stripping off the purified inhibitor with $0.1N\ H_3PO_4$. Samples were then immediately neutralized by adding $1/10\ volume\ 1M\ K_2HPO_4$.

Rosetta calculations. Rosetta energies of all designed structures were generated using the Slow Relax routine ⁴⁴. 1000 decoys were calculated for each design. PDB coordinates and energy parameters for the lowest energy decoy for each design are included as supplemental files.

Circular Dichroism (CD). CD measurements were performed in 100mM KPO₄, pH 7.2 with a Jasco spectropolarimeter, model J-1100 with a Peltier temperature controller. Quartz cells with path lengths of 0.1 cm and 1cm were used for protein concentrations of 3 and 30 μ M, respectively. The ellipticity results were expressed as mean residue ellipticity, [θ], deg cm² dmol⁻¹. Ellipticities at 222 nm were continuously monitored at a scanning rate of 0.5 deg/min. Reversibility of denaturation was confirmed by comparing the CD spectra at 20°C before melting and after heating to 100°C and cooling to 20°C.

Measuring HSA and IgG binding affinity. Affinity of proteins to HSA and IgG was determined by their retention on immobilized ligand. HSA and rabbit IgG were immobilized by reaction with NHS-activated Sepharose 4 Fast Flow (Cytiva) according to the manufacturer's

instructions. The concentration of immobilized HSA was 100 μ M. The concentration of immobilized IgG was 50 μ M (i.e. 100 μ M Fc binding sites). Generally 0.2mL of a 5 μ M solution of the test protein was injected onto a 5mL column at a flow rate of 0.5mL/min. Determination of binding affinity assumes that binding is in rapid equilibrium such that the elution volume is proportional to the fraction of test protein bound to 100 μ M of binding sites. Proteins that are completely retained after 20 column volumes (CV) are assessed to have $K_D \leq 1\mu$ M. Completely retained proteins are stripped from the column with 0.1N H_3PO_4 at the end of the run.

Measuring protease inhibition. Competitive inhibition constants (K_I) are measured by determining the $K_{M(apparent)}$ for the substrate QEEYSAM-AMC and RASProtease(I)⁴¹ in the presence of 100nM test protein.

NMR Spectroscopy. Isotope-labeled samples were prepared at 0.2-0.3 mM concentrations in 100 mM potassium phosphate buffer (pH 7.0) containing 5% D₂O. NMR spectra were collected on Bruker AVANCE III 600 and 900 MHz spectrometers fitted with Z-gradient 1 H/ 13 C/ 15 N triple resonance cryoprobes. Standard double and triple resonance experiments (HNCACB, CBCA(CO)NH, HNCO, HN(CA)CO, and HNHA) were utilized to determine main chain NMR assignments. Inter-proton distances were obtained from 3D 15 N-edited NOESY and 3D 13 C-edited NOESY spectra with a mixing time of 150 ms. NmrPipe 57 was used for data processing and analysis was done with Sparky 58 . Two-dimensional 14 H}- 15 N steady state heteronuclear NOE experiments were acquired with a 5 s relaxation delay between experiments. Standard deviations in heteronuclear NOEs were estimated based on the background noise level. Chemical shift perturbations between B₁ and S_{b1} were calculated using $\Delta\delta_{\text{total}} = ((W_{\text{H}}\Delta\delta_{\text{H}})^2 + (W_{\text{N}}\Delta\delta\text{N})^2)^{1/2}$, where W_{H} is 1, W_{N} is 0.2, and $\Delta\delta_{\text{H}}$ and $\Delta\delta_{\text{N}}$ represent 1 H and 15 N chemical shift changes, respectively. For PRE experiments on S_{b1}, single-site cysteine mutant samples were incubated with 10 equivalents of MTSL ((1-oxyl-2,2,5,5-tetramethylpyrroline-3-

methyl) methanethiosulfonate, Santa Cruz Biotechnology) at 25°C for 1 hour and completion of labeling was confirmed by MALDI mass spectrometry. Control samples were reduced with 10 equivalents of sodium ascorbate. Backbone amide peak intensities of the oxidized and reduced states were analyzed using Sparky. Three-dimensional structures were calculated with CS-Rosetta3.2 using experimental backbone 15 N, 1 H_N, 1 H α 13 C α , 13 C β , and 13 CO chemical shift restraints and were either validated by comparison with experimental backbone NOE patterns (A₁, B₁, B₄, S_{b1}) or directly employed interproton NOEs (S_{a1}, S_{b2}) or PREs (S_{b1}) as additional restraints. One thousand CS-Rosetta structures were calculated from which the 10 lowest energy structures were chosen. For S_{b3}, CS-Rosetta failed to converge to a unique low energy topology, producing an approximately even mixture of S- and B-type folds despite the chemical shifts and NOE pattern indicating an S-fold. In this case, CNS1.1 59 was employed to determine the structure as described previously 46 , including backbone dihedral restraints from chemical shift data using TALOS 60 . Protein structures were displayed and analyzed utilizing PROCHECK-NMR 61 , MOLMOL 62 and PyMol (Schrodinger) 45 .

Table 1: Structure statistics for A₁, B₁, and B₄.

	A ₁	B ₁	B ₄	
A. Experimental chemical shift inp	uts			
13 C $lpha$	55	56	56	
¹³ Cβ	51	50	53	
¹³ CO	54	55	53	
¹⁵ N	54	55	54	
¹ H _N	54	55	54	
1 H $lpha$	-	55	-	
B. RMSDs to the mean structure (Over all residues	(Å)			
Backbone atoms	1.73±0.47	0.86±0.22	0.85±0.34	
Heavy atoms Secondary structures ^a	2.27±0.53	1.37±0.29	1.43±0.45	
Backbone atoms	0.90±0.29	0.67±0.22	0.62±0.25	
Heavy atoms	1.55±0.42	1.12±0.25	1.33±0.41	
C. Measures of structure quality (9 Ramachandran distribution	%)			
Most favored	98.78±1.64	92.31±3.35	92.36±2.71	
Additionally allowed	1.22±1.64	7.69±3.36	7.64±2.71	
Generously allowed	0.00±0.00	0.00±0.00	0.00±0.00	
Disallowed	0.00±0.00	0.00±0.00	0.00±0.00	
D. Backbone RMSDs to the paren				
Over all residues	2.48	0.62	0.64	
Secondary structures <i>E. PDB/BMRB codes</i>	1.21	0.57	0.49	
PDBDEV	00000083	00000084	00000085	
BMRB	50907	50910	50909	

^a The secondary elements used were as follows: A_1 , residues 5-23, 27-35, 39-53; B_1 , residues 2-8, 13-19, 23-37, 43-46, 51-55; B_4 , residues: 2-8, 13-19, 23-37, 42-46, 51-55.

^b The parent structure for A_1 is G_A (PDB 2FS1). The parent structure for B_1 and B_4 is G_B (PDB 1PGA). RMSDs were calculated by superimposing the mean structure from the NMR ensemble with either the mean structure (in the case of 2FS1) or the X-ray structure (in the case of 1PGA) for the parent.

Table 2: Structure statistics for S_{a1} , S_{b1} , S_{b2} , and S_{b3} .

	S _{a1}	S _{b1}	S_{b2}	S _{b3}
A. Experimental restraint inputs				
NOE restraints				
Sequential (i-j=1)	89	-	35	40
Medium range (1< i-j≤ 5)	35	-	31	10
Long range (i-j > 5)	92	-	67	66
Hydrogen bond restraints	88	-	82	83
ΓALOS dihedral angle restraints	_	_	_	91
Total NOE restraint inputs	304		215	290
PRE restraints		41		
B. Experimental chemical shift inpu	ıts			
3 C $lpha$	88	79	83	_
³ Сβ	86	70	76	_
³ CO	69	67	70 72	_
⁵ N	88	69	72 76	_
H _N	88	69	76 76	-
Hlpha	76	45	70 61	-
$^{\sqcaplpha}$ C. RMSDs to the mean structure (A		45	01	-
Over all residues ^a	-1)			
Backbone atoms	2.77±0.82	5.47±1.86	2.34±0.60	2.46±0.64
Heavy atoms	3.51±0.85	6.32±1.86	3.00±0.62	3.29±0.61
Secondary structures ^b				
Backbone atoms	1.07±0.23	3.79±1.50	1.08±0.24	0.68±0.14
		$(0.71\pm0.23)^{c}$		
Heavy atoms	1.84±0.37	`4.37±1.42 [´]	1.78±0.32	1.42±0.25
·		$(1.24\pm0.30)^{c}$		
D. Measures of structure quality (%	6)	,		
Ramachandran distribution	,			
Most favored	86.46±4.07	92.35±2.38	92.19±1.83	86.30±2.38
Additionally allowed	13.54±4.07	7.54±2.60	7.81±1.83	10.52±2.68
Generously allowed	0.00±0.00	0.00±0.00	0.00±0.00	1.47±1.54
Disallowed	0.00±0.00	0.12±0.38	0.00±0.00	1.96±1.10
E. Backbone RMSDs to the parent				
Over all residues	2.37	0.49	6.16	11.67
Secondary structures	0.88	0.41	3.39	3.20
F. PDB/BMRB codes	0.00	0.11	0.00	0.20
	7MN1	7MQ4	7MN2	7MP7
PDB				

Data availability statement The NMR structures are available in the PDB (Accession Codes: 7MN1, 7MQ4, 7MN2, 7MP7) and PDBDEV (00000083, 00000084, 00000085). All other data are available from the corresponding authors on reasonable request.

^a Over all residues used as follows: S_{a1} , 1-95; S_{b1} , 4-85; S_{b2} , 1-93; S_{b3} , 1-87. ^b The secondary elements used are as follows: S_{a1} , residues 2-10, 16-32, 40-44, 59-67, 72-81, 86-92; S_{b1} , residues 5-12, 17-24, 27-41, 46-50, 55-59, 73-83; S_{b2} , residues 2-9, 23-32, 43-48, 59-65, 71-80, 85-91; S_{b3} , residues 4-10, 24-37, 40-46, 51-57, 62-71, 79-85.

 $[^]c$ RMSDs for S_{b1} minus the putative α 2 region: residues 5-12, 17-24, 27-41, 46-50, 55-59.

^d The parent structure for S_{a1}, S_{b2}, and S_{b3} is PDB 1RIS. The parent structure for S_{b1} is PDB 1PGA. In this case, the structure alignment is over the 56 amino acid B-region of S_{b1}.

Author contributions B.R., Y.H., and Yw.C. contributed equally. Protein design: Yw.C., B.R., E.C., J.O., P.B.; Performed thermodynamic and binding analyses: B.R., Yw.C., R.S., P.B.; Performed dynamic light scattering experiments: T.G.; Performed NMR experiments/structural analysis: Y.H., Yh.C., T.S., T.K., J.O.; Wrote paper: J.O. (NMR and structural analysis), Yw.C., B.R., P.B. (remaining sections).

Competing Interests statement The authors declare no competing interests.

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