Cysteine-free Rop: A four-helix bundle core mutant has wild-type stability and structure but dramatically different unfolding kinetics

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Abstract: Cysteine residues can complicate the folding and storage of proteins due to improper formation of disulfide bonds or oxidation of residues that are natively reduced. Wild-type Rop is a homodimeric four-helix bundle protein and an important model for protein design in the understanding of protein stability, structure and folding kinetics. In the native state, Rop has two buried, reduced cysteine residues in its core, but these are prone to oxidation in destabilized variants, particularly upon extended storage. To circumvent this problem, we designed and characterized a Cys-free variant of Rop, including solving the 2.3 Å X-ray crystal structure. We show that the C38A C52V variant has similar structure, stability and in vivo activity to wild-type Rop, but that it has dramatically faster unfolding kinetics like virtually every other mutant of Rop that has been characterized. This cysteine-free Rop has already proven useful for studies on solution topology and on the relationship of core mutations to stability. It also suggests a general strategy for removal of pairs of Cys residues in proteins, both to make them more experimentally tractable and to improve their storage properties for therapeutic or industrial purposes.

Keywords: Rop; four-helix bundle; cysteine-free; unfolding kinetics

Introduction
Protein engineering for improved physical properties, such as stability, folding kinetics or solubility, remains a difficult challenge.1 A major focus has been the repacking of the hydrophobic core of proteins, because a number of lines of evidence suggest that hydrophobic core packing is the key parameter affecting the stability of most proteins. These residues tend to be the most conserved for a particular protein family, and the most intolerant of mutation.2,3 Remarkable successes in packing or repacking protein cores have come from rational and computational design, as well as directed evolution.4–8 But there is still no accurate predictive model for the thermodynamic or kinetic consequences of even a single point mutation in the core of a protein.

Engineering cysteine residues into or out of a protein has been a major area of investigation. Disulfide bonds can dramatically stabilize structures, and have been successfully engineered for this purpose.9–11 It is also frequently desirable to engineer cysteine-free variants of proteins for study. Cysteines complicate the folding and storage of proteins,
both because disulfide bond formation tends to be a slow step in folding and because oxidation of cysteines (usually to adventitious disulfides) can result in misfolding and oligomerization. This is especially problematic for destabilized mutants of proteins. For example, a cysteine-free variant of T4 lysozyme was engineered by Matthews and colleagues as the basis for dozens of mutants designed to understand the structural and thermodynamic consequences of amino acid changes. A variant of human superoxide dismutase without free cysteines is resistant to thermal inactivation and has improved *E. coli* expression.

An interesting question in engineering cysteines is, to what (or from what) residues should one make a cysteine mutation? In active sites, a reasonable choice is serine, although serine is less nucleophilic and less acidic than cysteine. It is often underappreciated that cysteine is quite hydrophobic and much larger than serine due to the oxygen-to-sulfur substitution. In fact, cysteine is intermediate between alanine and valine both in volume and in the Kyte and Doolittle hydrophathy scale. Moreover, hydrogen bonds between the Cys thiol and backbone carbonyl oxygens’ would be expected to be much weaker than those commonly observed from the Ser or Thr alcohol, due to the poorer $pK_A$ match of the thiol proton. The BLOSUM62 matrix shows that the highest probability substitution throughout evolution of Cys is for Ala (log-odds of zero), followed by Ile, Leu, Met, Ser, Thr, and Val (log-odds of $-1$).

Coiled coils and helical bundles have proven especially useful for protein engineering and in elucidating the effects of mutations on protein stability. For example, the four-helix bundle Rop is a homodimer of 63 amino acid monomers arranged in an antiparallel fashion (Fig. 1). Extensive mutations have been made to the core and loop of Rop to probe their effects on stability, folding and structure. Recently, we have reported in vivo and in vitro screens for the folding and stability of Rop, with the goal of correlating the effects of many kinds of mutations with changes in thermodynamics. A proof-of-principle library of mutations in the central core of Rop produced a library of active variants with a range of physical properties (TJM and Lynne Regan, manuscript in preparation). Wild-type Rop has reduced cysteine residues buried in the core at positions 38 and 52, and less-stable and molten globular variants were found to oxidatively dimerize upon storage. These proved difficult to reduce, even upon boiling in SDS-PAGE loading buffer containing dithiotothreitol, presumably because once a disulfide forms between the monomers a second forms intramolecularly with little entropic penalty.

To circumvent this problem, we sought to engineer a cysteine-free Rop variant with native-like activity, structure and stability. To this end, we used our cell-based activity screen to examine combinations of Ser, Ala, and Val in positions 38 and 52. We found that a C38A C52V variant of Rop is stable and active, and the X-ray crystal structure demonstrates that there is little perturbation as a result of these mutations. However, Cys-free Rop unfolds much more rapidly than wild-type Rop, which has anomalously slow unfolding for a small protein lacking prolines or disulfide bonds. We have recently employed this useful variant as the basis for a comprehensive study of Rop core mutations using a dye-based screen for thermal stability we dubbed High-Throughput Thermal Scanning (HTTS).

**Results**

**Construction, activity, and stability**

From the crystal structure of wild-type Rop (1ROP), a hydrogen bond between the thiol of Cys38 and the backbone carbonyl oxygen of Thr19 appears possible (3.4 Å), but it is mostly surrounded by hydrophobic residues and appears to be well-packed (Fig. 2). The thiol of Cys52 is slightly further from the backbone carbonyl oxygen of Leu48 (3.5 Å), and it appears to be somewhat less well packed. Both positions 38 and 52 are in the a position of the canonical heptad repeat, which are typically small amino acids; however, the next-to-last layers of the Rop hydrophobic core are reversed, and position 52 would be expected to be a large amino acid (Fig. 1).

We decided to initially mutate Cys38 and Cys52 to

![Figure 1. The structure of wild-type Rop. Rop is an antiparallel homodimer of 63 amino acid monomers, placing the reduced Cys38 and Cys52 residues near each other across the dimer interface. At right, in the monomer, residues in the hydrophobic core of Rop are shown as spheres and labeled with their position (a, d, or e) in the heptad repeat. Note that in the last repeat, R55 (d) is mostly exposed and F56 (e) instead packs into the core. Rendered from 1ROP with PyMOL (Delano Scientific).](image)
Ser/Ser, Ala/Ala, and Val/Val. The genes for these variants were produced by PCR assembly of four ~62-mer synthetic oligonucleotides. The genes were cloned both into pACT7lac to assess function and into pMR10125 for overexpression.

Figure 3 shows the results of the in vivo functional assay. Rop controls the copy number of ColE1 plasmids in E. coli, and the basis of the screen is expression of green fluorescent protein driven from a ColE1 plasmid. In this implementation (the “positive screen”), active variants yield a high level of GFP expression, while an inactive variant shows a low level. The Ser/Ser mutant is not active. The other three constructs show a level of function equal to wild-type Rop. Analysis of ColE1 plasmid levels by miniprep and gel electrophoresis confirm this result (not shown).

Although both the Ala/Ala and Val/Val variants are active, only the Ala/Ala variant appeared to be expressed at significant levels under control of a T7 promoter. The reason for the lack of expression of Val/Val is not clear. The Ala/Ala variant has a similar mean residue ellipticity to wild-type Rop at 222 nm, indicating similar secondary structure. However, it is considerably less thermally stable than the Cys/Cys wild-type (T_m of 56°C vs. 70°C, Fig. 4).

Because Cys52 appears to be less well packed in the structure of wild-type Rop, we decided to construct and examine an Ala/Val mutant (C38A C52V). This variant is active by the cell-based screen. Its activity is also evident from DNA miniprep to examine plasmid levels, and it expresses at high levels from a T7 promoter (6–7 mg L^{-1} culture, not shown). Essentially all of the protein appears to be in the soluble fraction of the lysate. It can be concentrated to very high levels (>10 mg mL^{-1}) with no
signs of precipitation upon storage at 4°C for months.

The Ala/Val variant has similar helical content to wild-type Rop as judged from CD, and it is slightly more thermally stable than wild-type Rop (T_M of 74°C). While Ala/Val (D_{1/2} ~ 4.1M) is considerably more stable to urea (Fig. 4) than Ala/Ala (D_{1/2} ~ 2.3M), it is somewhat less stable than wild-type Rop (D_{1/2} ~ 4.6M). Because Rop is a homodimer, its folding is bimolecular, but its unfolding is unimolecular. Consequently, we perform all CD experiments at a defined standard state of 50 μM protein (with respect to the monomer). Taking the molecularity into account by the method of Dalal et al., we measure the stability of wild-type Rop to be 16 kcal mol⁻¹ (compared to 14.6 kcal mol⁻¹ reported by Dalal et al.) and C38V C52V to be 11 kcal mol⁻¹. The m-value changes from 2.1 kcal mol⁻¹ M⁻¹ for wild-type (Dalal et al. report 2.3 kcal mol⁻¹ M⁻¹) to 1.4 kcal mol⁻¹ M⁻¹ for Ala/Val. (See Table I for a summary of these parameters.)

We performed differential scanning calorimetry on Ala/Val Rop and compared the values to wild-type from the work of Munson et al. (Table II). The T_{1/2} values (temperature half-way through the calorimetric transition) at 0.3 mM are opposite those determined by CD spectroscopy at 50 μM (compare to Table I). Ala/Val Rop has a smaller enthalpic stabilization (ΔH_{cal}), but its ΔC_p value is within experimental error of the value for wild-type.

**Unfolding kinetics**

Wild-type Rop is particularly slow to unfold in chemical denaturant. Complete unfolding in the presence of 4M GdnHCl is reported to require more than a day. We examined the rate of unfolding of wild-type and Ala/Val Rop in urea. Even at high concentrations of urea, wild-type Rop requires more than 2 days to unfold, whereas C38A C52V requires less than an hour. By selecting several concentrations of urea where the proteins are more than 90% unfolded at equilibrium, it is possible to extrapolate the unfolding rates of the proteins in buffer (that is, no urea), assuming the logs of the rates are linearly related to the urea concentration over the full range (Fig. 5). The half-time (t_{1/2}) of unfolding of the Ala/Val mutant is 16 min, whereas the t_{1/2} for wild-type Rop is about 38 h, or ~1.5 days.

**Structure**

We solved the X-ray crystal structure of C38A C52V Rop to approximately 2.3 Å (Table III). The Ala/Val Rop crystallized in MES between pH 5.7–5.9 with 10% glycerol and 25–30% methanol as the

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**Table I. Thermodynamic and Kinetic Parameters of Rop Variants**

<table>
<thead>
<tr>
<th>Variant</th>
<th>M.R.E. (deg cm⁻² dmol⁻¹)</th>
<th>T_M (°C)</th>
<th>D_{1/2} (M urea)</th>
<th>m-value (kcal mol⁻¹ M⁻¹)</th>
<th>ΔG^H,O (kcal mol⁻¹)</th>
<th>k_u (s⁻¹)</th>
<th>t_{1/2} (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>w.t. (Cys/Cys)</td>
<td>21,000</td>
<td>70</td>
<td>4.6</td>
<td>2.1</td>
<td>16</td>
<td>5.1 x 10⁻⁶</td>
<td>2300</td>
</tr>
<tr>
<td>Ala/Val</td>
<td>20,000</td>
<td>74</td>
<td>4.1</td>
<td>1.3</td>
<td>11</td>
<td>7.3 x 10⁻⁴</td>
<td>16</td>
</tr>
<tr>
<td>Ala/Ala</td>
<td>20,000</td>
<td>56</td>
<td>2.3</td>
<td>0.9</td>
<td>n.d.</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
</tbody>
</table>

Equilibrium parameters were determined from 2 to 3 trials. Unfolding rates were determined from 2 or 3 independent trials at 4–7 different urea concentrations (Supporting Information).

* a M.R.E., mean residue ellipticity at 222 nm and 25°C; n.d., not determined.
precipitant. These conditions were suitably cryoprotective without additives. The crystal diffracted to 1.96 Å on Ohio State’s rotating-anode home source, but 93.3% completeness was only seen to 2.32 Å. All of the data were used in the refinement. The phases were estimated by molecular replacement with wild-type Rop (1ROP). The structure was rebuilt and refined, applying TLS refinement and refinement with riding hydrogens. The only notable geometric distortions in the fully-refined structure are the somewhat long bonds between N and C termini meet. At the N-terminus, the Thr2 ψ angle is rotated nearly 180°.

As with 1ROP, there is no clear density for the last six residues, which compose a charged “tail” that is poorly resolved due to its flexibility. There are three regions with modest differences from 1ROP, the first two at the open end of the hairpin where the N- and C-termini meet. At the N-terminus, the Thr2 ψ angle is rotated nearly 180° between the two variants. This is likely mostly because 1ROP begins with the sequence MTK... and Ala/Val Rop begins GTK... , as it was generated from a TEV protease cleavage before Gly1. Secondly, there is some perturbation at the C-terminus around the C52V mutation, most notably forcing the phenyl group of Phe56 toward the open end of the hairpin. There also appears to be some small deviation of the backbone in this region, but we see density for residue 57 which was not seen in 1ROP, so it is difficult to say if this is mostly a result of minor refinement differences. Thirdly, near the closed end of the hairpin, there is a small displacement of the backbone in the vicinity of Leu23 toward helix 2, possibly compensating for the volume loss in the C38A mutation. The Cα and Cβ positions of Ala and Leu overlay with the corresponding positions in the wild-type, and one of the Cγ atoms points along the same vector as Cγ-S in the wild-type at position 52. Other than the slight displacement of Phe56, there are essentially no changes to the side chains around either mutation, and even the Phe56 shift is due to a slight movement of the backbone and not a dramatic change in the χ angles. Other side chain differences in the rotameric states of surface Arg, Glu, and His residues are likely to arise from differences in refinement and/or multiple conformations.

**Discussion**

Rop has proven to be an important model protein for understanding the effects of mutations on structure, stability and folding kinetics, especially in the core

Table II. Differential Scanning Calorimetry

<table>
<thead>
<tr>
<th>Temperature ($^\circ$C)</th>
<th>∆H (kcal mol$^{-1}$)</th>
<th>∆Cp (kcal mol$^{-1}$ deg$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>w.t. (Cys/Cys)$^a$</td>
<td>73.9 ± 0.8</td>
<td>110 ± 6</td>
</tr>
<tr>
<td>Ala/Val$^b$</td>
<td>67.8 ± 0.8</td>
<td>65 ± 4</td>
</tr>
</tbody>
</table>

Numbers are ± standard deviation of three trials.

$^a$ From Munson et al.$^{19}$ in 100 mM phosphate, pH 7, 200 mM NaCl.

$^b$ 50 mM phosphate, pH 6.3, 300 mM NaCl.

Table III. X-Ray Data Collection and Refinement

<table>
<thead>
<tr>
<th>Data collection wavelength (Å)</th>
<th>Redundancy (r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.54</td>
<td>7.26</td>
</tr>
</tbody>
</table>

Resolution range (Å) 29.63–2.32 (2.46–2.32)

Space group C121

Cell dimensions $a$, $b$, $c$ (Å) 44.7, 40.3, 31.7

$\alpha$, $\beta$, $\gamma$ (°) 90.0, 102.0, 90.0

Mosaicity (°) 0.82

$R_{merge}$ 0.030 (0.079)

$I/\sigma I$ 47.4 (22.6)

Completeness (%) 93.3% (74.5%)

Redundancy 7.26 (7.04)

Refinement

Reflections (work$^c$/pairs/test) 4,885/2,675/242

$R_{work}/R_{free}$ 16.3/25.0

Average B-factor (Å$^2$) 26.2

Number of atoms

Protein 903

Solvent 69

R.M.S. deviations

Bond lengths (Å) 0.013

Bond angles (°) 1.088

$^a$ Reflections were observed to 1.96 Å, but the completeness beyond 2.32 Å is low. The data are 97.1% complete to 2.46 Å, and the overall completeness over the full resolution range is 66.1%.

$^b$ Number corresponds to 29.63–1.96 Å.

$^c$ Work reflections count the $F_{\text{obs}}$ (+) and $F_{\text{obs}}$ (–) reflections from the Bijvoet pairs separately.
and loop. As a homodimeric four-helix bundle, Rop is sufficiently simple to make rational mutations, and a great deal is known about the association of helices as coiled-coils. Still, Rop has afforded many surprises that could not easily be predicted, such as the topological inversion of a core variant repacked with Ala and Ile,\textsuperscript{27} or the topological rearrangement of the A31P mutant to a “bisecting U”.\textsuperscript{28} Recently, we introduced a means of measuring the melting temperatures of 96 variants of a protein at once (High-Throughput Thermal Scanning, or HTTS), and we demonstrated its use with Rop core variants.\textsuperscript{23} However, the buried Cys residues in the core at positions 38 and 52 complicate the mutagenesis of Rop, because destabilized mutants are prone to the formation of adventitious disulfides. This affects the reversibility of thermal denaturation and reduces the storage time of the protein. Consequently, we sought to generate a Cys-free variant with very similar structure and thermodynamic properties for further

Figure 6. X-ray crystal structure of Ala/Val Rop. (a) Overlay of the backbones of the symmetry-derived dimers of wild-type Rop (cyan carbons) and Ala/Val Rop (green carbons). The positions of the C38A and C52V mutations are shown as 50% transparent spheres for the wild-type Cys side chains. There is little change in the overall structure. (b) Overlay of the monomer backbones with core residues rendered as thick sticks. The backbone in helix 1 opposite C38A has moved slightly toward position 38. Near C52V, a small movement of the backbone has displaced F56 toward the open end of the hairpin. (c) Resides with atoms within 5 Å of the Cys38 sulfur are nearly unchanged except for the slight backbone shift toward the site of mutation. (d) It is evident that Val38 would have clashed with position of Phe56 in the wild-type, and it is displaced slightly by movement of the backbone. In (c) and (d), C38, C52, and F56 are shown in 50% transparent spheres with cyan carbons in the wild-type, and A38, V52, and F56 are shown in solid spheres with green carbons in Ala/Val. Rendered with PyMOL from 1ROP and the structure solved here, 3K79.
mutagenesis to understand the effects of mutation on protein stability. Cys-free variants of proteins such as T4 lysozyme and human superoxide dismutase have proven very useful in related studies.

Here we have shown that the C38A C52V variant of Rop is active, has wild-type like thermal stability, exhibits a cooperative thermal denaturation, and has similar although slightly reduced stability to urea denaturation. In contrast, the Ser/Ser variant was not active, the Val/Val variant did not express in appreciable quantities, and the Ala/Ala variant was substantially thermally destabilized. The decrease in the calorimetrically measured $\Delta H$ for folding suggests that slight imperfections in packing and bond angles, for example, have been introduced relative to wild-type Rop, and this is in accord with the slight perturbations observed around the two mutation sites in the crystal structure. The similar thermodynamic stability of wild-type Rop and the Ala/Val variant coupled with the lower $\Delta H$ suggest that the Ala/Val variant has considerably more favorable entropy of folding.

One way in which the Ala/Val variant does differ dramatically from wild-type Rop is in its much faster unfolding kinetics. The reason for Rop’s very slow unfolding is not clear; it contains no disulfide bonds or proline residues, it is a very small protein, and its variants with much faster unfolding kinetics (such as Ala/Val) are active, can be purified in large amounts, and do not aggregate. By way of comparison, several other core variants of Rop have extremely different folding kinetics. Rop can be considered to have eight 4-residue layers in its hydrophobic core. For wild-type, or in mutants wherein the central two or four layers are replaced with Ala2-Leu2, the unfolding rates were found to be around $10^{-4}$ s$^{-1}$ (extrapolated from GdnHCl denaturation); when six or eight layers are repacked, the rates increase to $10^{-2}$–$10^{0}$ s$^{-1}$.18 Cys38 is replaced in the four-layer variants and beyond, but Cys52 is not replaced until the six-layer variants. The effects of mutations on Cys52 alone may be worth exploration. Recently, Dalal et al. examined several homologs of Rop from other bacteria.26 Intriguingly, Cys38 is conserved among five identified homologs of Rop, but Cys52 is mutated to Ile or Leu in four of the five homologs. The two homologs examined experimentally both unfold more rapidly than Rop, one about 10-fold faster and one about $10^4$-fold faster. Both are C52L.

The Ala2-Leu2-6 (AL6) and Ala2-Ile2-6 (AI6) variants of Rop are both stable, but the AI6 variant does not bind RNA.8 When AI6 was crystallized, it was found in a parallel (or syn) topology, the opposite of that observed for wild-type Rop (anti, Fig. 7).27 The much faster unfolding of AL6 compared to wild-type Rop led Levy et al. to speculate that fast-unfolding variants might prefer the syn topology.29 In fact, we found that AI6 variant unfolds much faster than wild-type, but AL6 unfolds even faster, in the dead time of the CD experiment (J. E. Levy, unpublished). In collaboration with Deniz and Onuchic, we recently demonstrated by single-molecule FRET that at zero denaturant, both AL6 and AI6 adopt syn topology, while wild-type adopts anti.30 The AL6 variant can uniquely form both syn and anti at low denaturant. Both of these variants repack the Cys38 and Cys52 positions, and it will be interesting to examine the solution topological preferences of Ala/Val itself. It also remains to be seen if the differences in folding enthalpy and entropy are themselves related to the folding rates and solution topological preferences. It is notable that the Ala/Val variant was used as a labeling control in the smFRET experiments.

The unfolding rate we measured here for wild-type Rop is about an order of magnitude slower than Munson et al. measured, and the $T_M$ is about 6°C higher. This is probably mostly because of slight differences in the conditions: we used 50 μM protein, 50 mM phosphate, pH 6.3, 300 mM NaCl, and the previous study used 20 μM protein, 100 mM phosphate, pH 7, 200 mM NaCl. It is also possible that extrapolation back to zero denaturant from urea gives a different result from GdnHCl, perhaps because of the increasing ionic strength of the GdnHCl solutions.

Ala/Val Rop crystallized as an antiparallel homodimer, consistent with its activity. Overall, there is very little change to the structure as compared to wild-type. There is essentially no change in the positions of the residues known to be required

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Figure 7. Topologies of Rop variants. Wild-type Rop (1ROP, anti) and AI6 Rop (1F4M, syn) were rendered with PyMOL. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
for RNA binding: Asn10, Phe14, Gln18, or Lys25.31 Lys3 is weakly implicated in binding and is in a different rotameric state, but there was apparently not extensive density for this residue in 1ROP since the Cα and ε-amino groups are not shown. Very minimal perturbations are seen in the structure, principally in the backbone opposite C38A and with a small displacement of Phe56 next to C52V due to some backbone shifting.

Since buried cysteine residues are quite hydrophobic and have a side-chain volume that is intermediate between Ala and Val, we suggest that replacement of pairs of Cys residues with one Ala and one Val may be a good general strategy for cysteine mutagenesis. For example, we have even been able to replace a disulfide bond in another protein with Ala/Val, where either of the individual Cys → Ala mutants results in significant loss of activity (V. Shete & TJM, manuscript in preparation). This also suggests that it may be possible to introduce new disulfide bonds into proteins by mutating adjacent Ala and Val residues. The C38A C52V mutant of Rop has already proven extremely useful to us in development of a screen for protein stability. Since the unfolding kinetics of Ala/Val are more like virtually every core variant we have engineered and other small proteins lacking prolines and disulfides, we suggest this makes Ala/Val Rop a more useful reference state than wild-type for biophysical analysis. This Cys mutagenesis strategy may prove useful in many other contexts as well, such as removal of Cys residues to improve E. coli expression or to improve the storage properties of therapeutic proteins.

Materials and Methods

Cloning

Oligonucleotides (Sigma-Genosys) were used to assemble genes with point mutations at positions 38 and 52 of wild-type Rop (see Supporting Information Table S1). The mutation combinations were Ala/Ala, Ser/Ser, Val/Val, and Ala/Val. Each gene was amplified with two sets of primers to form “screening” genes that were used for the in vivo functional assay, and “expression” genes that contained N-terminal His6 and TEV sites. Screening genes (without His6 or TEV sites) were digested with NcoI and BamHI (New England Biolabs) and spin-column purified (Qiagen). Screening vector pACT7lac was digested with the same and gel-purified (Qiagen). Expression genes were digested with Bsal and NdeI and spin-column purified. Expression vector pMR101 was digested with the same and gel-purified. The digested genes were ligated (T4 DNA ligase (NEB)) with their respective vectors at 16°C, spin-column purified, and transformed into DH10B. Plasmids used for expression were transformed into BL21 (DE3) for protein purification.

Screening

The cell-based assay for Rop activity was performed as described.24 Briefly, genes in the screening vector were transformed into electrocompetent DH10B already containing the screening plasmid pUCBAD-GFPuv. Cultures were plated onto LB agar medium containing 100 μg mL⁻¹ ampicillin, 35 μg mL⁻¹ kanamycin, 0.1 mM IPTG, and 0.0005% arabinose (w/v), and incubated overnight at 42°C. Cellular fluorescence was visualized under hand-held UV illumination.

Protein purification

Rop variants in pMR vectors were overexpressed in BL21(DE3) in 1 L 2YT media. The culture was grown to log phase (OD600 ~0.7) and induced with 0.1 mM IPTG. The cells were harvested after growing at 30°C overnight or 37°C for 3 h. Frozen cell pellets were resuspended in 25 mL lysis buffer (50 mM Tris-HCl pH 8, 300 mM NaCl, 10 mM imidazole), mixed with 5 mM MgCl2, 0.5 mM CaCl2, 0.17 μg mL⁻¹ DNase I, 0.17 μg mL⁻¹ RNase A, 0.1% Triton X-100, and 0.3–1 mg mL⁻¹ HEW lysozyme, and allowed to incubate on ice for 30 min. The solution was then sonicated at 300 W for 3× 30 s with 2 min on in between pulses. After centrifugation, the cleared lysate was mixed on ice with 1 mL Ni-NTA agarose (Qiagen) for 1 h. The slurry was poured into a prefritted column (Bio-Rad), washed with wash buffer (lysis buffer with 20 mM imidazole), and the protein was eluted with elution buffer (lysis buffer with 250 mM imidazole). TEV protease (0.5 mg) was added twice to the protein and incubated each time overnight at RT or 3 h at 30°C. The solution was desalted with PD-10 columns (GE Amersham) and mixed with 1 mL Ni-NTA agarose and exchanged into appropriate buffer for subsequent analysis.

CD spectroscopy

Data were collected on an Aviv 202 Circular Dichroism Spectrometer. Rop variants were analyzed at 222 nm (using a calculated extinction coefficient of 1490 M⁻¹ cm⁻¹), in CD buffer (50 mM sodium phosphate, pH 6.3, 300 mM NaCl). Thermal denaturations were acquired at 1°C min⁻¹, 25–90°C, at 222 nm. Urea denaturations were performed in the same conditions but using varying concentrations of urea, with spectra acquired after equilibrating 24–48 h at RT, at 222 nm.

DSC

DSC measurements were made on a Microcal VP-DSC. Ala/Val Rop samples were in 500 μL with 2 mg mL⁻¹
(0.3 mM) protein in CD buffer. Denaturation was observed from 10 to 95°C at 1.5°C min⁻¹. Curves were fit to the data using Origin assuming a progressive baseline and a dissociative dimeric model (see below).

Data analysis
To calculate the $\Delta G^{H2O}$, the thermodynamic stability of the protein in buffer (i.e., zero denaturant), we followed the method of Dalal et al., assuming an equilibrium between the folded dimer and unfolded monomer.

$$N_2 = 2U$$  (1)

The equilibrium constants for each urea concentration $D$ were determined from fraction unfolded ($F_U$), fraction folded ($F_N$), and total protein concentration ($C_p$) using:

$$K_D = \frac{C_p F_U^2}{F_N}$$  (2)

These were converted to $\Delta G$ values from $-RT \ln K_D$ and the $\Delta G^{H2O}$ and m-values were determined by fitting a line:

$$\Delta G = \Delta G^{H2O} - m[D]$$  (3)

The rate constants for unfolding at each concentration of denaturant were determined by fitting single exponential functions ($A_k e^{-kt}$) to the time traces (Supporting Information Fig. S1). The rate constant for unfolding in water, $k_u$, was linearly extrapolated from the rate constants for folding at given concentrations of denaturant, $D$:

$$\log k_u = \log k_u^{H2O} + m_u[D]$$  (4)

Crystal growth and data collection
Rop was purified as above, followed by gel filtration purification using a Superdex 75 10/300 GL column (GE Life Sciences). The protein was then exchanged into a mild buffer (10 mM PIPES pH 6.3, 50 mM NaCl) and concentrated, if necessary, to approximately 2 mg mL⁻¹ as determined by Bradford assay. Sitting drop and hanging drop trials were performed. Plates were obtained with both ammonium sulfate and 2-methyl-2,4-pentanediol (MPD) precipitant. Rhombohedral prisms were obtained from hanging drop wells using 1 mM reservoirs containing 25–30% methanol, 100 mM MES, pH 5.7–5.9, 300 mM NaCl, and 10% glycerol. Crystals appeared overnight at 22°C and grew for 3–5 days. Diffraction data were collected on a Rigaku R-AXIS IV++ at −160°C and a detector distance of 200 mm. Images were collected at 0.5° increments with exposure times of 2 min. The images were processed, scaled, and reduced using the program d*TREK.32

Structure determination and refinement
Molecular replacement modeling was performed with Phaser,33 using 1ROP as the model, and ARP/wARP34 was used to rebuild the structure. Refinement was performed using phenix.refine.35 The structure was hand-corrected with Coot.36 Later refinement included TLS refinement (using TLSMD37 parameters), followed by a final refinement using hydrogens added with Molprobity.38

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References