Solution structure and biochemical characterization of a spare part protein that restores activity to an oxygen-damaged glycyl radical enzyme

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Received: 30 April 2019 / Accepted: 21 June 2019 / Published online: 27 June 2019
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Abstract
Glycyl radical enzymes (GREs) utilize a glycyl radical cofactor to carry out a diverse array of chemically challenging enzymatic reactions in anaerobic bacteria. Although the glycyl radical is a powerful catalyst, it is also oxygen sensitive such that oxygen exposure causes cleavage of the GRE at the site of the radical. This oxygen sensitivity presents a challenge to facultative anaerobes dwelling in areas prone to oxygen exposure. Once GREs are irreversibly oxygen damaged, cells either need to make new GREs or somehow repair the damaged one. One particular GRE, pyruvate formate lyase (PFL), can be repaired through the binding of a 14.3 kDa protein, termed YfiD, which is constitutively expressed in E. coli. Herein, we have solved a solution structure of this ‘spare part’ protein using nuclear magnetic resonance spectroscopy. These data, coupled with data from circular dichroism, indicate that YfiD has an inherently flexible N-terminal region (residues 1–60) that is followed by a C-terminal region (residues 72–127) that has high similarity to the glycyl radical domain of PFL. Reconstitution of PFL activity requires that YfiD binds within the core of the PFL barrel fold; however, modeling suggests that oxygen-damaged, i.e. cleaved, PFL cannot fully accommodate YfiD. We further report that a PFL variant that mimics the oxygen-damaged enzyme is highly susceptible to proteolysis, yielding additionally truncated forms of PFL. One such PFL variant of ~77 kDa makes an ideal scaffold for the accommodation of YfiD. A molecular model for the rescue of PFL activity by YfiD is presented.

Electronic supplementary material
The online version of this article (https://doi.org/10.1007/s00775-019-01681-2) contains supplementary material, which is available to authorized users.

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Introduction

Radical-based chemistry allows enzymes to catalyze an impressive array of molecular transformations such as those that involve cleavage and formation of C–C bonds (Fig. 1a) [1–5]. This chemistry comes at a price, however, as a radical species must be generated on the enzyme, a process that usually involves a metallocofactor [6]. The radical species must also be stored between turnovers, controlled during turnover, and protected from or repaired after oxygen damage. Some radical-generating cofactors have native protective mechanisms; adenosylcobalamin (AdoCbl) typically generates its highly reactive 5′-deoxyadenosyl (5′-dAdo) radical through Co–C bond homolysis only when substrate is present, and then reforms the Co–C bond when each turnover is complete, limiting both unwanted chemistry and interception of the radical species [3]. Although this strategy is relatively successful, AdoCbl is an expensive cofactor, requiring upwards of thirty enzymes for its biosynthesis [7, 8]. In contrast, glycyl radical enzymes (GREs) have an inexpensive cofactor, a glycyl radical species (i.e. a glycine that is missing a hydrogen atom), which transiently forms the catalytically essential thiyl radical species during turnover (Fig. 1a). A glycyl radical species does require a metallocofactor-containing activase for its formation [9, 10], but a single activase can post-translationally modify many of its associated GREs,
and the modification need only happen once, as the glycyl radical is reformed from the thiyl radical on every round of turnover (Fig. 1a). Additionally, the activases are S-adenosylmethionine (AdoMet) radical enzymes [9], which have been referred to as ‘poor man AdoCbl enzymes’ since they use inexpensive cofactors [4Fe–4S] and AdoMet for radical generation [2, 11–13]. GREs are thus an efficient option for radical-based enzymology, but unlike AdoCbl-dependent enzymes, they are very sensitive to inactivation by molecular oxygen. In these enzymes, oxygen-induced cleavage occurs at the glycyl radical site, splitting the enzyme into two pieces. For strictly anoxic-living microbes, GREs are an excellent option, powerful catalysts that are inexpensive for the cell to produce; however, for microbes that live under microaerobic conditions, GREs can be an acceptable option if damage repair is a possibility. Here, we investigate a novel repair mechanism for the central microbial metabolic GRE, pyruvate formate lyase (PFL).

PFL cleaves the C–C bond of pyruvate to produce formate and acetyl-CoA, providing acetyl-CoA to metabolic pathways including the citric acid cycle during anaerobic glycolysis (Fig. 1a) [14, 15]. Although the glycyl radical of PFL is oxygen sensitive, the enzyme is constitutively expressed, albeit with about tenfold higher PFL expression under anoxic conditions [9, 16, 17]. Under transient microaerobic environments, *Escherichia coli* upregulate expression of the *yfd* gene product YfiD [18, 19], a small protein that appears to be able to rescue the enzymatic activity of oxygen-damaged PFL [20]. Key regulators of anaerobiosis and pH in *E. coli*, *arcA*, *fnr* and *fur*, appear to be responsible for YfiD upregulation [21–24]. YfiD has 127 amino acids, the last sixty of which share 77% sequence identity to the

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**Fig. 1** PFL reaction, activation, and rescue by YfiD. **a** PFL catalyzes conversion of pyruvate and CoA to acetyl-CoA and formate. The glycyl radical (G734) of PFL is generated through a direct hydrogen atom abstraction by PFL activating enzyme, or PFL-AE, a member of the AdoMet radical enzyme superfamily. Catalysis occurs through a putative catalytic thiyl radical. **b** PFL-AE (orange) installs a glycyl radical on PFL in a reaction that is thought to require a conformational change of a region of PFL known as the glycyl radical domain (GRD) (maroon). Upon oxygen exposure, the C-terminus of PFL is cleaved at the glycyl radical site (star). YfiD (blue), which is also capable of being activated by PFL-AE, is able to bind to PFL and restore PFL activity.
The last sixty residues of *E. coli* PFL, the portion of PFL that contains both the glycyl radical (Gly734) and a region of the enzyme (residues 713–759), referred to as the glycyl radical domain (GRD) (Fig. 2) [25]. The GRD was previously shown by circular dichroism (CD) spectroscopy to undergo a conformational change when PFL binds to its activating enzyme (PFL-AE), presumably to expose the buried Gly734 for radical installation (Fig. 1b) [26]. It is also this region (734–759) of the activated protein that is cleaved upon oxygen exposure. The ability of a highly homologous small protein to restore activity to a much larger enzyme suggests that YfiD is the biological equivalent of a 'spare part' (Fig. 1b) [20].

The molecular mechanism by which YfiD is able to restore activity is unknown. Previous work established that PFL-AE can install a glycyl radical on YfiD [20] and we assume that activated YfiD must bind to cleaved PFL (cPFL) to restore activity, but exactly how this restoration takes place is unclear. The cleavage that occurs cuts the loop containing Gly734 in half. This glycyl radical loop is positioned next to the loop that contains the transient thiyl radical species in the core of PFL’s 10-stranded β-barrel (Fig. 2a, b). YfiD is much longer, at 127 residues, than the cleaved portion of PFL (734–759), and it is uncertain if more residues of cPFL will be displaced as a result of YfiD binding or if only a small fraction of YfiD will bind to cPFL. A sequence alignment of PFL GRD and YfiD shows that PFL Gly734 is directly aligned with YfiD Gly102. The residues in this region of the alignment are completely conserved for a stretch of 22 amino acids. Furthermore, the sequence similarity between YfiD and PFL is restricted to the last 60 residues, raising the question of whether YfiD will adopt a PFL-like fold in the non-conserved segments (Fig. 2c).

Although the exact number of residues involved in binding remains unknown, YfiD must be bound tightly enough to cPFL that the inserted glycyl radical is stable. To investigate these questions, we have taken a structural approach using nuclear magnetic resonance (NMR) spectroscopy.

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**Fig. 2**: Structure of PFL (PDB ID: 2PFL) monomer and sequence alignment of the C-terminal region of PFL and YfiD. **a** Crystal structure of PFL showing residues 633–711 in blue and residues 712–759 that comprise the glycyl radical domain (GRD) in red. The thyl radical loop is in yellow, and the rest of the PFL monomer is in gray. **b** Topographical representation of PFL structure. **c** Sequence of YfiD and PFL residues 633–759, showing the secondary structural elements of PFL below the sequences.
and circular dichroism (CD) spectroscopy to determine the solution structure of the E. coli YfiD protein and explore its thermal stability. We have further created a truncated form of PFL that mimics the oxygen-cleaved enzyme (cPFL) and have investigated whether cellular components cause additional proteolysis of cPFL in the presence of YfiD. Our results taken together allow us to propose a model for how YfiD binds and restores activity to cPFL.

**Results**

**YfiD is a partially structured/partially unstructured protein**

Representative spectra of YfiD from 2D $^1$H–$^{15}$N heteronuclear single quantum coherence (HSQC) experiments show good peak separation indicative of a folded protein (Fig. 3a). To solve the structure, standard triple-resonance experiments were used for backbone and side-chain assignments (see Materials and Methods), and distance restraints based on nuclear Overhauser effects (NOEs) were obtained from $^1$H–$^{15}$N-NOESY-HSQC and $^1$H–$^{13}$C-NOESY-HSQC experiments.

The calculations of the YfiD NMR structure converge well, as indicated by the ensemble of the ten lowest energy structures shown in Fig. 3b. However, the solved structure only contains the C-terminal half of YfiD (residues 68–127). Very few cross peaks could be identified for the N-terminal region of YfiD (residues 1–65), suggesting that this region is disordered, as depicted by the dotted line. c Sequence and secondary structure of YfiD (cyan) and residues 633–679 of PFL (colored as in Fig. 2)
The last 62 residues of YfiD form an autonomous glycyl radical cofactor

Given that the oxygen-cleavage site on PFL is in the middle of the Gly radical loop, it is not surprising that YfiD duplicates the full Gly radical loop. A glycine residue at the N-terminus of a protein would likely not be able to form a stable glycyl radical species [29], even if it were a substrate for PFL-AE. Thus, it was expected that YfiD would contain an intact Gly radical loop, including the PFL C-terminal α10β helix that is cleaved, but it was unclear how much more of the PFL structure would be duplicated in YfiD. Although the high sequence homology between YfiD and PFL starts

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The first 60 residues of YfiD dramatically affect YfiD’s thermal denaturation

To investigate the thermal stability of a protein that appears to be partially unstructured, we employed far-UV circular dichroism (CD) spectroscopy. We also created a truncated form of YfiD (truncYfiD) in which the majority of residues found to be unstructured by NMR (residues 1–60) were deleted. Thermal denaturation measurements were carried out on both YfiD and truncYfiD to determine the protein melting temperatures under two different buffering conditions. The denaturation curves for YfiD reveal that the protein does not undergo a denaturation typical of a well-folded protein, as illustrated by the non-sigmoidal shape to the denaturation curve (Fig. 4a, b). This unusual unfolding behavior is consistent with a protein that has a large unstructured region, where that unstructured region dominates the CD spectrum [27, 28]. As a control, we tested full-length YfiD expressed in the presence of PFL to determine if co-expression leads to a better ordered YfiD protein; it did not. An analogous CD spectrum was obtained (Fig. S1). However, when the unstructured part of YfiD is genetically removed, a classic sigmoidal shape for the denaturation curve is observed (Fig. 4c, d). In other words, truncYfiD (residues 61–127) behaves as well-ordered, folded protein. Again, this result is consistent with the NMR structure, in which residues 68–127 are structured.

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Fig. 4 Circular dichroism spectroscopy thermal denaturation measurements for YfiD in the presence and absence of the first 60 residues. a, b The thermal denaturation curves for intact YfiD in two different buffers (MES, pH 6.5 and HEPES, pH 7.2). For the YfiD in HEPES buffer, a melting temperature cannot be determined as the algorithm does not converge. c, d The thermal denaturation curves for a YfiD variant that is missing the first 60 residues of the protein (truncYfiD) in two buffers (MES, pH 6.5 and HEPES, pH 7.2). Tm values are listed in panels.
at the beginning of PFL’s β10 strand, the residues preceding this strand could have formed similar secondary structure with a different sequence, but that does not appear to be the case. We do find, however, a full duplication of the PFL β10 strand and α10a helix, some 30 residues more than the part of PFL that is cleaved by oxygen. The inclusion of these ‘extra’ residues could serve multiple functions, including: providing binding affinity to PFL-AE; protecting the Gly radical during transfer from PFL-AE to cPFL; providing binding affinity to cPFL; and securing the Gly radical loop in the active site barrel for proper radical transfer to the Cys loop to initiate catalysis. In terms of binding to either PFL-AE for activation or PFL for restoration of enzyme activity, the use of a highly identical sequence in this structural unit of YfiD (a BLAST alignment of YfiD and PFL yields 77% identity for YfiD residues 64–127 aligned to PFL residues 696–759) allows YfiD to swap in for PFL residues with shape and charge complementary intact.

To further investigate how YfiD might interact with PFL-AE, we manually docked YfiD onto the X-ray structure of PFL-AE, emulating the proposed mode of binding of PFL to PFL-AE (Fig. 5) [25]. We observe that the two helices and the Gly radical loop of YfiD fit well to PFL-AE; however, YfiD’s y-β1 strand is not accommodated in this docking mode (Fig. 5).

**Proteolysis of PFL follows oxygen-induced cleavage**

The discovery that YfiD is structurally homologous to 60 residues of the C-terminus of PFL, rather than just to the 25 residues of PFL (734–759) that are cleaved upon oxidative damage, indicates that more residues of PFL might need to be displaced from the active site for YfiD to bind in the location that places the Gly radical loop correctly for chemistry to occur. Given that one cleavage event often leads to protein unfolding, which can cause previously buried cleavage sites to be exposed, we wanted to investigate whether cPFL is further unfolded by examining its susceptibility to proteolysis by cellular proteases. We co-expressed His-tagged YfiD and a truncated version of PFL that mimics the oxygen-damaged enzyme (cPFL) and purified the resulting complex by immobilized metal affinity chromatography (IMAC). In addition to full-length cPFL (81 kDa), we observe bands for two truncation products (Fig. 6), a more intense band that is consistent with a 77 kDa fragment and a less intense band that is consistent with a 71 kDa fragment. Cleavage occurring between residues 690–695 (t1PFL) and residues 635–640 (t2PFL) would be expected to generate fragments of 77 kDa and 71 kDa, respectively. A common E. coli proteolytic enzyme, chymotrypsin-like protease I, has sequence specificity consistent with cleavage in these regions. In fact, the ExPASy PeptideCutter prediction algorithm predicts two cleavage sites between residues 690 and 695 with >73% probability of being cleaved by E. coli chymotrypsin-like protease I, and one cleavage site between residues 635 and 640 with 58% probability of being cleaved by this common E. coli protease (Fig. 6b). These results are consistent with cPFL undergoing further protein unfolding events following oxygen-induced cleavage at Gly734, which exposes previously buried proteolysis sites.

**A single proteolysis event following oxygen-induced cleavage of PFL creates a PFL variant optimal for YfiD docking**

As described above, the YfiD structure duplicates all the Gly radical domain (red in Fig. 7a), as well as β10 strand (blue in Fig. 7a) and α10a helix (red in Fig. 7a), but does not duplicate β9 strand and α8b helix (blue in Fig. 7a, YfiD teal in Fig. 7b). The active site in cPFL would still be relatively buried (Fig. 7c) with no room for YfiD to bind. A single proteolysis event at the β9-α9 loop (Fig. 7a) would result in loss of β10 strand and α10a helix as well as the remaining residues of the GRD. A space filling illustration of such a PFL (t1PFL) shows an open active site that is ready for YfiD binding (Fig. 7d). Modeling YfiD into t1PFL shows an
impressive fit with very little need for structural rearrangement (Fig. 7d, e). When YfiD is positioned as shown in 7e, the Gly radical loop is located suitably for radical transfer to the thiyl radical loop, and y-β1 strand is situated such that it serves as a strand to complete the 10-stranded barrel of PFL (Fig. 7b).

**Discussion**

Since the discovery of YfiD by Knappe and coworkers in 2001 [20], the idea that enzymes have ‘spare parts’ has been a source of fascination. Here, we present the NMR structure of YfiD and consider the molecular basis by which this ‘spare part protein’ may function in the repair of the major microbial metabolic enzyme PFL. First and foremost, radical chemistry requires precise positioning of cofactors, residues, and substrates. Thus, the restoration of catalysis in PFL requires that an activated YfiD be precisely positioned in the active site of PFL such that the glycyl radical species is stable, allowing for reversible radical transfer to Cys418 and multiple turnovers. Prior to this work, it was not clear how much of PFL would be duplicated in the YfiD structure and whether the binding of the YfiD structural unit to cPFL would require conformational rearrangements. Although a structure of YfiD:PFL has not yet been obtained despite considerable effort, we do not believe that the observed proteolysis of cPFL is a coincidence. Due to the fact that the major truncated form of cPFL (t1PFL) is a species that contains an opening that is perfectly sized for YfiD (Fig. 7d, e), we now propose that oxygen-induced cleavage to form cPFL is followed by proteolysis to form t1PFL and that t1PFL is the enzyme variant to which YfiD binds (Fig. 8, steps 1–4).

Of course, YfiD binding cannot reconstitute PFL activity if YfiD does not contain a glycyl radical species. Thus, YfiD requires post-translational modification by PFL-AE to install a radical at position Gly102 [20] before YfiD assumes its catalytically competent position inside the PFL barrel structure. Although it has been shown that YfiD is a substrate for PFL-AE [20], little is known about the nature of the complex that is formed. In general, our understanding of glycyl radical formation is limited [1]. Currently, the only structural data we have on how a glycyl radical domain is activated by an activase are from the crystal structure of a 7-mer peptide substrate bound to PFL-AE [25]. We do not know how much of the PFL glycyl radical domain binds to PFL-AE to afford activation nor do we know the exact binding mode of PFL to PFL-AE.

Previously, we presented a model for the binding of residues 712–759 of PFL to PFL-AE that was based on the shape and charge complementarity [25]. If we now superimpose the NMR structure of YfiD onto this PFL:PFL-AE docked structure, we observe that the two helices and the Gly radical loop of YfiD fit well, but the y-β1 strand does not (Fig. 5). It is possible that the docked PFL:PFL-AE structure is not an accurate model for how YfiD binds to PFL-AE; another possibility is that YfiD undergoes a conformational rearrangement of the y-β1 strand before binding to PFL-AE, potentially assisted by the binding of YfiD to t1PFL in a pre-activation step (Fig. 8, steps 3–4).

Although biochemical data have not yet been obtained to demonstrate that YfiD, t1PFL, and PFL-AE form a ternary complex.

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**Fig. 6** cPFL is subject to proteolysis. a SDS-PAGE of cPFL. Molecular weights of cPFL, 81 kDa, and two additional truncation products at 77 kDa (t1PFL) and 71 kDa (t2PFL) were extrapolated using a 10–200 kDa protein standard. These molecular weight bands correspond to the expected length of cPFL ending at residue 733, PFL truncated between residues 690–695 (t1PFL), and PFL truncated between 635 and 640 (t2PFL), respectively. b Crystal structure of the PFL monomer showing residues 734–759 (end of C-terminus) in maroon, residues 695–733 in blue, the catalytic Cys loop in purple, and the rest of the monomer in gray. Upon exposure to oxygen, PFL is cleaved at Gly734. This cleavage exposes two truncation sites for *E. coli* chymotrypsin-like protease I activity. Positions of cleavage sites are indicated with stars.
complex prior to activation, the formation of such a complex would explain how two potential pitfalls could be avoided. First, there must be a mechanism that protects the glycyl radical species on YfiD from being quenched before YfiD is positioned into the protective PFL active site. Formation of a ternary complex could provide this protection. Second, it would not be in the best interest of the cell to generate a glycyl radical species on YfiD if there was no damaged PFL to repair. Requiring that complex formation with a damaged PFL be the first step in the YfiD activation process ensures that YfiD is not activated when it is not needed. In this sense, YfiD’s y-β1 can be thought of as a safety lever, preventing radical formation by PFL-AE until a damaged PFL ‘pulls’ the lever down (Fig. 5). With these ideas in mind, we have revised our proposed mechanism of PFL rescue by YfiD (Fig. 8, Electronic Supplementary Material Video 1). In particular, we now show both the unstructured region and the y-β1 strand of YfiD bound to t1PFL with the unstructured region affording general binding affinity and the y-β1 strand replacing the lost β10, completing the 10-stranded barrel fold. Collectively, these parts of YfiD play a functional role in ‘sensing’ PFL damage and in tethering YfiD to t1PFL prior to YfiD activation by PFL-AE.
A revised model for YfiD rescue. When exposed to oxygen, activated PFL is cleaved at its glycyl radical site, G734 (cPFL). The cleaved region of PFL (G734 through the C-terminus) is depicted in red. In order for YfiD to fit into the pocket, additional residues that include residues of β10 strand of PFL (blue) must be lost, forming a tPFL variant. From here, we propose that the y-β1 strand of YfiD completes the 10-stranded barrel of PFL and the disordered N-terminus of YfiD helps secure this spare part protein to tPFL. After YfiD is anchored to PFL, a conformational change occurs in which the glycyl radical domain (GRD) flips out, allowing PFL-AE to activate YfiD. Once activated, the GRD of YfiD enters the open PFL active site, resulting in a reactivated PFL:YfiD complex. This complex should remain stable for multiple rounds of turnover with YfiD acting as a noncovalent subunit.

**Materials and methods**

**Plasmid construction**

We received YfiD in a p-CAL-n-EK vector as a gift from the Broderick lab. To prepare the truncated YfiD (truncYfiD) construct used in the circular dichroism experiments, the Q5® Site-Directed Mutagenesis Kit (New England Biolabs) protocol was used to delete the first 180 base pairs from the *yfid* gene (corresponding to a deletion of 60 amino acids at the N-terminus), in the p-CAL-n-EK vector. The following primers were used for this deletion: forward primer, 5′-GTG AAA CCA GAA GTT CGC-3′, and reverse primer, 5′-CAT ATG TAT ATC TCC TTC TTA AAG-3′. After the deletion was performed, a hexahistidine-TEV cleavage site was added at the 5′ end of truncYfiD, again using the NEB Q5® Site-Directed Mutagenesis Kit. All primers were designed using NEBaseChanger™. All mutagenesis experiments were confirmed through Sanger sequencing by GENEWIZ, Inc.

To prepare the pPFL:YfiD co-expression plasmid used in the SDS-PAGE experiment, the *yfid* gene was inserted into the second MCS of pCOLA-Duet using NdeI and XhoI cut sites, and a hexahistidine-TEV cleavage site was added at the 5′ end of this gene (Genscript). Next, a synthesized 733-PFL (mimicking cleaved PFL, or cPFL) insert (Genscript) was mutated to add a PciI cut site at the 5′ end and a SalI cut site at the 3′ end and inserted into the first MCS of pCOLA-duet at the NcoI and SalI cut sites (NcoI and PciI have compatible sticky ends).
**Protein expression and purification**

YfiD, truncYfiD, and cPFL:YfiD plasmids were all transformed into T7 Express cells (New England Biolabs) for expression. The following expression protocol was used for all three constructs: samples were prepared in growth media supplemented with 100 μg/mL ampicillin (Affymetrix). Unlabeled YfiD, truncYfiD, and cPFL:YfiD were expressed in Luria-Bertani broth. Starter cultures were inoculated with single colonies and incubated at 37 °C overnight at 220 RPM. 10 mL of starter culture was used to inoculate 1 L of media, which was incubated at 37 °C at 220 RPM until the optical density at 600 nm was 0.6. Expression was induced by addition of 1 mM IPTG (RPI Corp.), followed by an additional 3 h of incubation at 37 °C. Cells were spun down, and cell paste was frozen and stored at −80 °C prior to purification.

Uniformly 15N labeled YfiD (U–15N-YfiD) and uniformly 13C15N labeled YfiD (U–13C15N-YfiD) for NMR experiments were expressed in M9 minimal media supplemented with 10 mM magnesium sulfate (EMD-Millipore) and 0.03 g/L thiamine hydrochloride (Sigma-Aldrich). 1 g/L 15N-ammonium chloride (Cambridge Isotopes Laboratories) and 25 mL of 40% glucose were added to provide 15N and C for U–15N-YfiD. 1 g/L 15N-ammonium chloride (Cambridge Isotopes Laboratories) and 3 g/L 13C-glucose (Cambridge Isotopes Laboratories) were added to provide 15N and 13C for U–13C15N-YfiD. Note that the deuterium from the 13C-glucose exchanges rapidly in the non-deuterated media. Aside from growth media, the expression protocol was the same as described above for unlabeled YfiD.

To purify U–15N-YfiD and U–13C15N-YfiD, cell paste was thawed and lysed at 4 °C while nutating for 1 h with 10 mL lysis buffer for each 5 g of cell paste (lysis buffer: 20 mM HEPES (Fisher Bioreagents) pH 7.2, 10 mM magnesium chloride (EMD-Millipore), 5% W/V glycerol (VWR Analytics), 1% V/V Triton X-100 (Sigma-Aldrich), 1 cOmplete EDTA-free tablet (Roche) and 1 μL Benzonase Nuclease (EMD-Millipore). The resulting suspension was sonicated with ¼” tip at 60% power for 4×2 min cycles of 2 s on and 2 s off. Lysate was clarified at 18,000 RPM, and the protein was eluted from a DEAE column (GE Healthcare) in a linear gradient from 0 to 500 mM imidazole. 20 mM HEPES, pH 7.2. Fractions containing truncYfiD were buffer exchanged into imidazole-free 20 mM HEPES, pH 7.2. truncYfiD was treated with hexahistidine-tagged TEV protease at a 1:100 (protease to truncYfiD) concentration ratio and incubated overnight at 4 °C. TEV protease was separated from untagged truncYfiD by collecting the flow-through and low imidazole fractions from a cOmplete His-Tag purification column (Roche). Protein was concentrated in a 10 kDa Amicon spin concentrator (EMD-Millipore).

cPFL:YfiD cell paste was thawed and lysed following the same protocol as described for U–15N-YfiD and U–13C15N-YfiD. Lysate was clarified at 18,000 RPM. The cPFL:YfiD complex, with a hexahistidine-tag on the N-terminus of YfiD, was purified from a His SpinTrap column (GE Healthcare Life Sciences) by increasing the concentration of imidazole from 0 to 500 mM imidazole, with 20 mM NaCl, 20 mM HEPES, pH 7.2. Fractions of cPFL:YfiD eluted at 100 mM imidazole were pooled.

Protein concentration was determined by absorbance at 280 nm using the molar extinction coefficients of 11,460 M⁻¹ cm⁻¹ for YfiD, 5960 M⁻¹ cm⁻¹ for truncYfiD, and 49,565 M⁻¹ cm⁻¹ for cPFL:YfiD.

**SDS-PAGE**

An SDS-PAGE gel was run to distinguish the different cleavage products of cPFL. SDS-PAGE was analyzed using a 7.5% Mini-PROTEAN® TGX™ precast gel (Biorad). Unstained Protein Standard, Broad Range (10–200 kDa, New England Biolabs) was used as the molecular weight standard. Although cPFL was purified with YfiD, YfiD had to be run off the gel in order to observe sufficient band separation of the cPFL cleavage products.

**Circular dichroism**

Similar far-UV CD spectra were recorded on an Aviv Model 202 CD Spectrometer for experiments found in Fig. 4 and a JASCO Model J-1500 Circular Dichroism Spectrometer for experiments shown in Fig. S1. Both spectrometers are equipped with a jacketed cell holder connected to a circulating water bath, and 0.1-cm path length cuvettes were used (Starna). Protein samples were prepared of 80 μM YfiD or 80 μM truncYfiD in 10 mM HEPES, pH 7.2 and 10 mM MES, pH 6.5 (Sigma-Aldrich). The thermal denaturation...
experiments recorded the CD signal at 220 nm at 5 °C intervals with a 30 s equilibration time at each temperature range, over the temperature range 20–90 °C. Thermal denaturation curves were obtained by plotting the CD signal at 220 nm as a function of temperature and the melting temperature was determined by fitting the curves to the following two-state model equation:

\[
y = \frac{(y_f + m_f T) + (y_u + m_u T) \exp(\Delta H_m/R(1/T_u-1/T))}{1 + \exp(\Delta H_m/R(1/T_u-1/T))}
\]

where \( y \) is the measured ellipticity, \( R = 8.3145 \text{ J K}^{-1} \text{ mol}^{-1} \), \( y_f \) and \( y_u \) are the intercepts of the pre- and post-transition baselines, \( m_f \) and \( m_u \) are the slopes of the pre- and post-transition baselines, \( T \) is the temperature (K), \( \Delta H_m \) is the enthalpy at the unfolding transition and \( T_m \) is the melting temperature (K). The equation is fit in R Statistical Software [35] using nonlinear least squares to fit all parameters.

**NMR spectroscopy**

NMR spectra of 0.7 mM U–15N-YfiD (18 mM HEPES pH 7.2, 2.7 mM (NH₄)₂SO₄ 10% V/V D₂O) were collected on CMR 600 E, a 600 MHz NMR spectrometer at Massachusetts Institute of Technology’s Francis Bitter Magnet Lab. NMR spectra of 1.3 mM U–13C₁⁵N-YfiD (18 mM HEPES pH 7.2, 2.7 mM (NH₄)₂SO₄ 10% V/V D₂O) were collected on Varian Inova 800 MHz spectrometer, equipped with a cryogenic triple-resonance probe, at University of Connecticut Health Center NMR facility. Chemical shifts were referenced to water. All NMR data were processed using NMRPipe [36] and analyzed using NMRFAM-SPARKY [37]. Standard triple-resonance experiments were used for backbone and side-chain assignments, including 2D ¹H–¹⁵N HSQC and ³¹C–¹⁵N HSQC, and 3D HNCO, HNCA, HNCACB, CBCACONH, HCCHTOCSY, HBHACONH, and ¹⁵N resolved ¹H–¹H TOCSY. Distance restraints based on nuclear Overhauser effect (NOE) were obtained from ¹H–¹⁵N-NOESY-HSQC and ³¹H–¹⁵C-NOESY-HSQC. HSQC spectra were obtained between each experiment to assess quality of the protein. Peak picking and NOE assignments were performed using NMRFAM-SPARKY [37], CCPN [38], and CYANA [39]. Structure calculations were performed using NOE distances and dihedral angle constraints, and refined with molecular dynamics. The backbone RMSD from the medoid lowest energy structure is 0.07 Å for the well-defined core residues of the protein, indicating convergence of the NMR structure solution. NMR assignments and details have been deposited to the Biological Magnetic Resonance Bank (ID 30611) and the 10 lowest energy conformers have been deposited to the PDB (ID 6OWR).

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