Steric trapping reveals a cooperativity network in the intramembrane protease GlpG.
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Abstract

Membrane proteins are assembled through balanced interactions among protein, lipids and water. Studying their folding while maintaining the native lipid environment is necessary but challenging. Here we present methods for analyzing key elements in membrane protein folding including thermodynamic stability, compactness of the unfolded state and folding cooperativity under native conditions. The methods are based on steric trapping which couples unfolding of a doubly-biotinylated protein to binding of monovalent streptavidin (mSA). We further advanced this technology for general application by developing versatile biotin probes possessing spectroscopic reporters that are sensitized by mSA binding or protein unfolding. By applying these methods to an intramembrane protease GlpG of Escherichia coli, we elucidated a widely unraveled unfolded state, subglobal unfolding of the region encompassing the active site, and a network of cooperative and localized interactions to maintain the stability. These findings provide crucial insights into the folding energy landscape of membrane proteins.

Understanding the free energy landscape of protein folding requires determination of the free energy levels and conformations of states populated during folding as well as analysis of energy barriers to reach the native conformation1. Experimentally, this task has been carried out by equilibrium and kinetic folding studies using denaturants that can readily shift
population distribution between folded and unfolded states. However, the overall shape of the folding energy landscape substantially changes in the presence of denaturants, and certain short-lived higher energy states may not be detected in denaturing conditions. Thus, studying protein folding under native conditions is necessary for a full survey of the folding energy landscape. For water-soluble proteins, methods such as hydrogen-deuterium exchange (HDX), NMR relaxation dispersion and proteolysis have revealed the dynamic and multi-state nature of the native conformational ensemble which is critical to protein function. However, such features remain largely unexplored for membrane proteins. The poor accessibility of solvent water to the interior of micelles and bilayers and the large sizes of protein-micellar and protein-liposomal complexes have made it difficult to apply these methods for characterizing the native ensemble of membrane proteins.

Steric trapping is a promising tool for investigating thermodynamic stability and folding of membrane proteins directly under native conditions. The method couples unfolding of a target protein labeled with two biotin tags to competitive binding of bulky monovalent streptavidin molecules (mSA, MW=52 kD) (Fig. 1a for detailed description).

Although promising, it is yet difficult to apply steric trapping to various types of membrane proteins. The method requires two essential features: two site-specifically conjugated biotin labels on a target protein and a probe to monitor mSA binding or protein unfolding. Site-specific biotinylation has been achieved by labeling of engineered cysteine residues with thiol-reactive biotin derivatives. For detection of unfolding, widely-used tools such as tryptophan fluorescence and circular dichroism cannot be used due to large signal interferences from mSA molecules. A method for direct detection of mSA binding has not been developed yet. Thus, the application has been limited to proteins possessing convenient unfolding readouts such as absorbance of a conformation-sensitive intrinsic chromophore (e.g. retinal in bacteriorhodopsin) and enzymatic activities (e.g. dihydrofolate reductase and diacylglycerol kinase). In this study, we developed a generalized steric trapping strategy which utilizes novel thiol-reactive biotin probes containing spectroscopic reporter groups for sensitive detection of mSA binding and protein unfolding. This strategy was applied to analyze the thermodynamic stability, compactness of the unfolded state and folding cooperativity of the six-helical bundle intramembrane protease GlpG of E. coli.

GlpG is a member of the rhomboid protease family widely conserved in all kingdoms of life. Rhomboid proteases are involved in diverse biological processes by activating membrane-bound signaling proteins or enzymes via cleavage of a specific peptide bond near the membrane. Due to the functional importance of rhomboid proteases and detailed structural information available (28 PDB entries, http://www.rcsb.org/), GlpG has emerged as an important model for studying the folding of helical membrane proteins. Critical regions for the stability have been identified using heat and sodium dodecylsulfate (SDS) denaturation tests of 151 variants. A kinetic folding study using SDS as a denaturant has suggested the existence of a compact folding nucleus in the folding transition state. A single-molecule tweezers study has shown that GlpG largely unfolds cooperatively at a constant tension.
Here, using steric trapping, we provide new insights into the folding energy landscape of GlpG in the absence of heat, chemical denaturants or pulling force. We elucidated an expanded heterogeneous conformational ensemble of the unfolded state, a structural region that undergoes subglobal unfolding, and an intricate network of cooperative and localized interactions to maintain the stability of GlpG.

RESULTS

Design and synthesis of new steric-trapping probes

Our steric-trapping probes are highlighted by three features that are integrated into one molecular tag (Fig. 1b): (1) a biotin group for binding mSA, (2) a thiol-reactive group for conjugation to engineered cysteine residues on a target protein, and (3) a fluorescent or paramagnetic reporter group whose spectroscopic signal is sensitized by mSA binding or protein unfolding. Each probe was synthesized by stepwise substitutions of building blocks possessing characteristic features to a lysine or cysteine template (Supplementary Results, Supplementary Notes). BtnPyr-IA (1) is a pyrene-based fluorescent sensor to detect mSA binding. When used to doubly label a target protein, pyrene fluorescence is remarkably sensitive to binding of quencher-labeled mSA by Förster resonance energy transfer (FRET). BtnRG-TP (2) is a paramagnetic sensor possessing a 1-oxyl-2,2,5,5-tetramethylpyrroline spin label to detect protein unfolding. The spin labels allow distance measurements in the native and steric-trapped unfolded state using double electron-electron resonance spectroscopy (DEER).

Steric trapping controls reversible folding of GlpG

To prove the principle of our steric trapping strategy employing the new probes, we used GlpG as a model and its proteolytic activity as a folding indicator. Here all studies were performed in dodecylmaltoside (DDM) micelles, in which a majority of functional and folding studies of GlpG have been carried out, and with the isolated transmembrane (TM) domain (residues 87–276) for which all structures of GlpG have been solved. For precise and efficient measurement of GlpG activity with membrane-bound substrates, we developed a fluorescence-based assay that can be transformed into a high-throughput format (Supplementary Fig. 1). The activity of the TM domain of GlpG was indistinguishable from that of the full-length protein (Supplementary Fig. 2).

For steric trapping, we first identified optimal residue pairs for cysteine substitution to conjugate thiol-reactive biotin labels. After testing of multiple single- and double-cysteine variants, two double-cysteine variants, P95C/G172C and G172C/V267C were selected (Fig. 2a). The biotin pair conjugated to P95C/G172C is located in the approximate N-terminal half of GlpG (95/172_N hereinafter) while the biotin pair to G172C/V267C is located in the C-terminal half (172/267_C). Individual single-cysteine variants P95C, G172C and V267C labeled with fluorescent BtnPyr-IA maintained the wild-type activity level (Fig. 2b, top) and this activity level was not significantly altered after binding of wild-type mSA (mSA-WT). Michaelis-Menten analysis of the proteolytic activity showed that $K_M$, $k_{cat}$ and $k_{cat}/K_M$ of the single-biotin variants bound with mSA were indistinguishable from those of unbound forms demonstrating that binding of one mSA molecule to each biotin site did not perturb
the structure and function of GlpG (Supplementary Fig. 2). The wild-type activity level was also maintained after labeling of double-cysteine variants. In marked contrast, saturated binding of mSA to two biotin labels on each variant induced a substantial loss of activity implying GlpG was trapped in the unfolded state (Fig. 2b, bottom).

Next, we tested if the steric-trapped unfolded state can refold after dissociation of bound mSA. Wild-type mSA binds biotin with an enormously high affinity ($K_{d,\text{biotin}} \approx 10^{-14}$ M) and slow dissociation rate ($k_{\text{off}} \approx 30$ days). Thus, we used mSA-S27A variant with a weaker biotin affinity (Supplementary Fig. 3) to facilitate dissociation of bound mSA by addition of excess free biotin (Fig. 1a). Both double-biotin variants which were inactivated with mSA-S27A significantly regained the activity upon addition of free biotin (Fig. 2b, bottom). For 95/172$_N$-BtnPyr$_2$, 50–70% of activity was regained while >90% was regained for 172/267$_C$-BtnPyr$_2$. Thus, we achieved reversible folding of GlpG without using denaturants by steric trapping.

**Steric-trapped unfolded state is widely unraveled**

So far, protein unfolding by steric trapping has been tested by the loss of enzymatic activity, decrease of retinal absorbance, or increased susceptibility to proteolysis. Although those features indicate unfolding, a possibility remains that the protein conformation trapped with mSA molecules is only locally distorted or still compact with significant residual tertiary interactions. Therefore, to elucidate the conformation of the steric-trapped unfolded state as well as to gain insights into the unfolded state ensemble of membrane proteins under non-denaturing conditions, we used a thiol-reactive biotin derivative possessing a spin label (BtnRG-TP) (Fig. 1b). By labeling of double-cysteine variants of GlpG with this probe, we have the advantage of both trapping the unfolded state and measuring the distances between spin labels using DEER. DEER allows for measurements of long-range (15–60 Å) inter-spin distances. It provides not only the most probable distance but also distance distribution, which is of great interest in characterization of the unfolded state. Here, we obtained inter-spin distances for 95/172$_N$-BtnRG$_2$ and 172/267$_C$-BtnRG$_2$ in their native, SDS-induced unfolded and steric-trapped unfolded states (Fig. 3). In the native states, the distance distributions between BtnRG labels were overall similar to those between well-characterized R1 spin labels (Supplementary Fig. 4) demonstrating that our BtnRG label is capable of distance mapping of protein conformation.

For both variants, SDS induced substantial broadening of the inter-spin distance distribution over the range from the native-like distances (15–35 Å) up to ~60 Å (Fig. 3, right panels), which indicates a heterogeneous conformational ensemble of the unfolded state in SDS. Interestingly, in non-denaturing DDM micelles, the steric-trapped unfolded states also exhibited similarly broad inter-spin distance distributions. The increase of the most probable distance from ~25 Å in the native state to ~55 Å in the steric-trapped unfolded state corresponds to a ~30 Å expansion of each half of the polypeptide chain covered by the respective biotin pair. This increased dimension is comparable to the whole diameter of native GlpG. Thus, our DEER data for GlpG rules out a compact unfolded state under non-denaturing conditions, which has been observed for several water-soluble proteins. We note that, because of the detection limit of DEER, even longer distance components (>60 Å)
may have existed but not been detected. Addition of dithiothreitol to break the disulfide bond between GlpG and the biotin label bound with mSA led to a regain of activity (>70%) indicating that a majority of the unfolded conformations were able to refold (Supplementary Fig. 5).

Steric repulsion between bound mSA molecules may have biased the conformational ensemble of the unfolded state. However, during the selection of optimal biotin pairs, we found that saturated binding of mSA to the biotin pairs conjugated to G94C/G172C and G172C/N271C, whose Cα-Cα distances were similar to those of 95/172N and 172/267C, completely retained the activity and therefore did not induce unfolding (Supplementary Fig. 6). This result implies that bound mSA molecules are allowed to coexist within close distances probably also in the steric-trapped unfolded state. Therefore, steric repulsion may not fully explain the expanded unfolded state.

We further characterized the conformational features of the steric-trapped unfolded state using proteolysis by chymotrypsin, which primarily targets aromatic residues prevalent throughout GlpG (Supplementary Fig. 7). While the unfolded state bound with two mSA molecules was gradually proteolyzed over ~30 min, either native GlpG or GlpG bound with one mSA molecule was not significantly proteolyzed. As a control, we tested proteolysis of casein which dominantly exists in random coil conformation in aqueous solution. Casein was rapidly proteolyzed within 1 min. Thus, we speculate that the steric-trapped unfolded state was significantly protected by secondary structures and micelles, but possessed more dynamic features than the native state.

DEER and proteolysis results demonstrate that steric trapping induced a true unfolded state, which was an ensemble of expanded dynamic and heterogeneous conformations. This work also represents the first measurement of the physical dimension of a helical membrane protein in its unfolded state under non-denaturing conditions.

**Stability of GlpG determined by steric trapping**

To develop a general steric-trapping strategy that does not depend on specific characteristics of a target protein, ideally the spectroscopic signal from the reporter group in our probe (Fig. 1b) should sensitively change upon either mSA binding or protein unfolding. Here we achieved a highly sensitive detection of mSA binding by employing FRET between pyrene on BtnPyr label and non-fluorescent quencher DABCYL on mSA (mSA_DAB) (Fig. 4a).

SDS denaturation and linear extrapolation of the denaturation data to zero-SDS mole fraction yielded the same stability (ΔG_U,SDS) for 95/172N-BtnPyr2 (8.4±1.5 kcal/mol) and 172/267C-BtnPyr2 (8.7±1.2 kcal/mol) (Fig. 4b and Supplementary Fig. 8), which was similar to that of the full-length wild type (8.2±1.4 kcal/mol). This result indicates that the two double-biotin variants possessed the same global stability as wild type GlpG. By design, steric trapping specifically captures transient unfolding of native interactions between a biotin pair. Thus, probing the stability with two biotin pairs located in different regions (Fig. 2a) provides a novel opportunity to test the folding cooperativity of GlpG.
Using FRET, we obtained high-quality binding isotherms between each double-biotin variant and three mSA\(_{\text{DAB}}\) variants with different biotin affinities (Fig. 4c). mSA\(_{\text{DAB}}\) binds the biotin label at each site with a similar affinity (Supplementary Fig. 3a). An essential element of steric trapping to determine protein stability is choosing an mSA variant whose binding to a biotin label (\(\Delta G_{\text{Bind}}\)) optimally competes with folding (\(\Delta G_{\text{U}}\)) to yield attenuated second binding in a desirable [mSA] range (Fig. 4a). Among mSA variants tested, mSA\(_{\text{DAB}}\)-S27A yielded an optimal separation of the first tight and second weaker binding phases (Fig. 4c). Parallel activity measurements showed that, for each GlpG variant, the second binding coincided with the activity loss (i.e. unfolding), which validated the unfolding-binding coupling. The same coupling was observed with a high-affinity variant, mSA\(_{\text{DAB}}\)-WT, further confirming that the activity loss strictly depended on the second binding of mSA\(_{\text{DAB}}\) (Supplementary Fig. 9).

Fitting of the second binding phases yielded the thermodynamic stability (\(\Delta G_{\text{U,ST}}, \text{ST: steric trapping}\), 5.8±0.2 kcal/mol for 95/172\_N-BtnPyr\(_2\) and 4.7±0.1 kcal/mol for 172/267\_C-BtnPyr\(_2\) (equation (4) in Online Methods) in non-denaturing DDM micelles (Fig. 4c). Both \(\Delta G_{\text{U,ST}}\)'s were significantly lower than the extrapolated stability from SDS denaturation (8.4–8.7 kcal/mol) but higher than the stability in a bicelle (6.5 \(k_B T\) equivalent to ~4 kcal/mol) extrapolated to zero force from single-molecule tweezers study. If GlpG unfolded cooperatively, the same \(\Delta G_{\text{U,ST}}\) would be expected regardless of the position of the biotin pair. However, while SDS denaturation yielded the same global stability for the two double-biotin variants, their stabilities obtained by steric trapping were comparable but significantly different by 1.1±0.2 kcal/mol.

**Subglobal unfolding of GlpG near the active site**

To track down the origin of the discrepancy between the stability obtained by steric trapping (\(\Delta G_{\text{U,ST}}\)) under a non-denaturing condition and the extrapolated stability obtained by SDS denaturation (\(\Delta G_{\text{U,SDS}}\)), we directly measured the stability of the two double-biotin variants using steric trapping in the range of SDS mole fraction (\(X_{\text{SDS}}=0–0.4\)), where a major fraction of GlpG existed in the folded state (folded fraction >0.9) (Fig. 4b). The \(\Delta G_{\text{U,ST}} \text{ vs } X_{\text{SDS}}\) plot (Fig. 5) revealed two major features that clearly deviated from the behavior predicted from linear extrapolation of SDS denaturation data. First, rather than following a linearly-decreasing trend, \(\Delta G_{\text{U,ST}}\) of both variants exhibited an upward curvature as \(X_{\text{SDS}}\) increased. Second, while \(\Delta G_{\text{U,ST}}\) of 95/172\_N-BtnPyr\(_2\) were overall larger than that of 172/267\_C-BtnPyr\(_2\), they remarkably converged at \(X_{\text{SDS}} \approx 0.4\) where the main unfolding transition by SDS began, and this convergence was maintained up to \(X_{\text{SDS}}=0.5\). This result confirms that the two variants possessed the same global stability.

The overall nonlinearity of \(\Delta G_{\text{U,ST}}\) against \(X_{\text{SDS}}\) implies a complex interaction between GlpG and DDM/SDS micelles. A similar disagreement between steric trapping and SDS denaturation has been reported for bacteriorhodopsin in DMPC/CHAPSO/SDS bicelles. In the case of GlpG, \(\Delta G_{\text{U,ST}}\) of both variants reached a maximum at \(X_{\text{SDS}} \approx 0.2\) but linearly decreased at higher \(X_{\text{SDS}}\) (Fig. 5). Notably, in the range of \(X_{\text{SDS}}=0.2–0.4\), the \(m\)-value of 95/172\_N-BtnPyr\(_2\) (14±2 kcal/mol/\(X_{\text{SDS}}\)), which represents the slope of \(\Delta G_{\text{U,ST}}\) against...
$X_{SDS}$ was significantly larger than that of 172/267$_C$-BtnPyr$_2$ (8±1 kcal/mol/$X_{SDS}$) but similar to those obtained by SDS denaturation (16–17 kcal/mol/$X_{SDS}$).

For water-soluble proteins, the $m$-value is correlated with the hydrophobic surface area exposed upon unfolding. Although the physical meaning of the $m$-value in SDS denaturation is still under debate, it is most likely related to the difference in the affinity of SDS for different states of the protein, hence to the degree of exposure of buried stabilizing interactions upon unfolding. Therefore, from the different denaturant sensitivities of the two double-biotin variants, we conclude that trapping of the unfolded state with the biotin pair 95/172$_N$-BtnPyr$_2$ led to substantial exposure of the buried surfaces throughout the protein, while trapping with the biotin pair 172/267$_C$-BtnPyr$_2$ mainly occurred through subglobal unfolding which exposed less buried-surface area. Steric trapping of 172/267$_C$-BtnPyr$_2$ detects transient separation between TM3 and TM6 to which biotin labels are conjugated, and TM6 contains a biotin label (V267C-BtnPyr) as well as His254 of the catalytic dyad (Fig. 6a). Thus, subglobal unfolding should directly involve disruption of the active site.

Subglobal unfolding has been frequently observed from HDX studies of water-soluble proteins, but not been reported for membrane proteins. Besides the different $m$-values, subglobal unfolding of GlpG is further supported by the lower stability of 172/267$_C$-BtnPyr$_2$ from steric trapping (Fig. 4c) and the reproducibly higher refolding yield of 172/267$_C$-BtnPyr$_2$ ($>90\%$) than that of 95/172$_N$-BtnPyr$_2$ (50–70\%) (Fig. 2b), implying different unfolded states.

We also note that $\Delta G^\text{U,ST}_{1}$ was larger than $\Delta G^\text{U,SDS}_{1}$ after it crossed the extrapolation lines at $X_{SDS}=0.1$ and this discrepancy became increasingly pronounced up to 2.8 kcal/mol at $X_{SDS}=0.4$ (Fig. 5). We reason that the larger $\Delta G^\text{U,ST}_{1}$ was primarily due to the conformational difference between the steric-trapped unfolded state and the SDS-induced unfolded state. Our DEER result supports this argument (Fig. 3). The steric-trapped unfolded state on average exhibited larger inter-spin distances than the SDS-induced unfolded state. Thus, steric trapping appears to induce more unraveled conformations than SDS at least for the interactions between the biotinylated sites. However, we are cautious with this direct comparison because the compactness of the SDS-induced unfolded state may change as a function of $X_{SDS}$ due to the effects of SDS on the size and shape of mixed micelles.

**Strategy to identify cooperative interactions**

The higher stability and more substantial unfolding obtained with 95/172$_N$-BtnPyr$_2$ indicates that the native interactions between this biotin pair in the N-terminal region are critical to the conformational integrity of the whole protein. On the other hand, the lower stability and subglobal unfolding obtained with 172/267$_C$-BtnPyr$_2$ indicates that the C-terminal region possesses differential folding properties from the N-terminal region. This result suggests complex energetic coupling between different regions in GlpG. To clarify this complexity, we developed a method to identify cooperative and localized interactions that contribute to the protein stability at a side-chain resolution (Fig. 6a).
First, we dissected GlpG into two subdomains: (1) the more stable N-terminal subdomain I encompassing TM1-L1-TM2-TM3-L3198 (ending at residue 198 in the L3 loop) whose unfolding was trapped with 95/172N-BtnPyr2, and (2) the less stable C-terminal subdomain II consisting of L3199-TM4-TM5-L5-TM6 (starting from residue 199) whose subglobal unfolding was trapped with 172/267C-BtnPyr2 (Supplementary Fig. 10 for dissection procedures). The uncertainty of the division point was ±20–30 residues.

Second, we made a single mutation (typically to alanine) in either subdomain to perturb a specific side-chain interaction in the background of 95/172N-BtnPyr2 and 172/267C-BtnPyr2. We referred to these background double-biotin variants as “wild type (WT)” because the wild-type native interactions were equally preserved in both as shown by SDS denaturation (Fig. 4b). We referred to two double-biotin variants possessing the same mutation as “mutants (Mut)”. Next, we probed the stability changes induced by the mutation with two different biotin pairs using steric trapping. We quantified the differential effect of the same mutation on the stability of each subdomain (ΔΔΔG) using equation (1) containing the stabilities of four variants:

\[
\Delta \Delta \Delta G^0_{U} = \left[ \Delta G^0_{U,95/172N-BtnPyr2} (WT) - \Delta G^0_{U,95/172N-BtnPyr2} (Mut) \right] - \left[ \Delta G^0_{U,172/267C-BtnPyr2} (WT) - \Delta G^0_{U,172/267C-BtnPyr2} (Mut) \right]
\]

\[
= \Delta \Delta G^0_{U,95/172N-BtnPyr2} (WT-Mut) - \Delta \Delta G^0_{U,172/267C-BtnPyr2} (WT-Mut)
\]

(1)

ΔΔG^0_{U,95/172N-BtnPyr2}(WT-Mut) and ΔΔG^0_{U,172/267C-BtnPyr2}(WT-Mut) designate the stability changes caused by the same mutation in the backgrounds of 95/172N-BtnPyr2 and 172/267C-BtnPyr2, respectively. Thus, ΔΔΔG^0_{U} represents the difference in the stability changes that are probed with two different biotin pairs upon the same mutation.

If a mutation causes a similar degree of destabilization for both double-biotin variants with a difference within thermal fluctuation energy (|ΔΔΔG^0_{U}| ≤ RT=0.6 kcal/mol; R: gas constant; T=298K), the mutated site engages in a “cooperative” interaction. That is, the perturbation by the mutation similarly propagates to both subdomains. Among the cases where |ΔΔΔG^0_{U}| > RT, if a mutation preferentially destabilizes the subdomain containing it, the perturbed interactions are “localized” within that subdomain. If mutation of a residue, which makes its side-chain contacts only with the subdomain containing it, preferentially destabilizes the other subdomain, we classified the perturbation as “over-propagated”.

Cooperativity network in GlpG

We targeted 20 residues covering key packing regions23 and analyzed their roles in the folding cooperativity of GlpG (Supplementary Table 1). The stability changes upon mutation ΔΔG^0_{U}(WT-Mut) obtained by steric trapping were reasonably well correlated with the changes in melting temperature ΔTm(WT-Mut)23, which validated our approach (Supplementary Fig. 11). 20 ΔΔΔG^0_{U} values were distributed over a wide range from −1.8 to 2.0 kcal/mol and their individual errors ranged from ±0.1 to ±0.4 kcal/mol, smaller than
We applied four cut-off values, \( \Delta \Delta G_{U} = -2RT, -RT, RT \) and \( 2RT \) (i.e. five sets of the cooperativity profile) to account for the wide distribution of \( \Delta \Delta G_{U} \) as well as to more precisely resolve the degree of cooperativity of each side-chain interaction.

We mapped the effects of mutations onto the structure, which we called the “cooperativity map” (Fig. 6b). Surprisingly, we observed significant clustering of cooperative and localized interactions in defined regions in the GlpG structure and divided their spatial distributions into four distinct groups. First, cooperative interactions of five residues, Met100, Leu161, Leu174, Thr178 and Ser201, clustered in the buried region which was surrounded by subdomain I and the subdomain interface near the center of the membrane. This cooperative cluster overlapped with one of the key packing regions previously identified\(^{23}\) and partially with the folding nucleus formed between TM1 and TM2 in the folding transition state\(^{24}\).

Second, all tested residues located in the folded L1 loop (Tyr138, Thr140 and Leu143) and the residue packed against L1 (Cys104) in subdomain I engaged in moderately \((RT \leq \Delta \Delta G_{U} \leq 2RT)\) or highly \((2RT < \Delta \Delta G_{U})\) localized interactions in subdomain I. This region is known to form non-native interactions in the folding transition state\(^{24}\). Third, Leu225 \((\Delta \Delta G_{U} < -2RT)\) and Gln226 \((-2RT \leq \Delta \Delta G_{U} < -RT)\) in TM5 in subdomain II, which were located at the subdomain interface and exposed to the water-micelle interface, respectively, were both classified as localized in subdomain II. TM5 is not tightly packed against the rest of the protein and does not significantly contribute to the thermostability\(^{23}\).

In the fourth cluster, interestingly, mutation of residues at the TM4/TM6 interface (Ala253, Gly261, Ala265 and Asp268) in subdomain II preferentially destabilized subdomain I, not the subdomain containing them. Particularly, Gly261 and Ala265 make their side-chain contacts entirely with the residues in subdomain II, but perturbing these interactions exerted larger impacts on the stability of subdomain I. Thus, we classified these residues as over-propagated. The TM4/TM6 interface harbors the catalytic dyad and plays a pivotal role in both stability and function of GlpG\(^{23}\). Especially, Gly261 and the dyad are absolutely conserved among rhomboid proteases\(^{44}\). Our result suggests that these conserved residues are also critical to the energetic coupling between different regions of GlpG. Breakage of the interactions near the C-terminus and its propagation towards the N-terminus is known to be the primary mechanism of the force-induced unfolding of GlpG\(^{25}\).

It should be noted that 5 among 20 tested mutations completely inactivated GlpG (Supplementary Table 1). Thus, our steric trapping strategy allowed stability measurements of not only functional but also non-functional variants, which had been difficult under the original steric-trapping framework. An ANOVA test of the five sets of \( \Delta \Delta G_{U} \) including the upper and lower standard-deviation limits of each \( \Delta \Delta G_{U} \) indicates that differences among these sets were statistically significant (Supplementary Fig. 12). While the two double-biotin variants bearing the same mutation exhibited differential stability change in DDM micelles, they possessed the same global stability from SDS denaturation (Supplementary Fig. 13). Therefore, we conclude that the networked side-chain interactions revealed in this work is a novel phenomenon that occurs under native conditions.
DISCUSSION

Here we presented a new steric-trapping strategy to investigate thermodynamic stability of membrane proteins and conformation of their unfolded state under native conditions by employing novel thiol-reactive biotin tags. Fluorescent BtnPyr allowed determination of the thermodynamic stability of GlpG through high-quality binding isotherms obtained by FRET. Paramagnetic BtnRG enabled characterization of the unfolded state based on the distance measurements using DEER. Because this combined strategy is not limited by either target-specific unfolding readout or specific lipid environments, it is applicable to other types of membrane proteins including nonfunctional and misfolded variants whose folding characterization is difficult under native conditions.

Unfolded state of proteins has gained significant interest because it determines thermodynamic stability with the folded state, directs folding mechanisms, and serves as a target for chaperoning and degradation\(^{45}\). However, conformation of the unfolded state of membrane proteins was difficult to study under native conditions due to its transient nature preventing biophysical analysis. By combining DEER and steric trapping, we elucidated a largely-unraveled dynamic and heterogeneous conformational ensemble of the unfolded state of GlpG in non-denaturing micellar solution. It is still an open question to what extent trapping would affect the protein conformation beyond the region containing the biotin pair. Investigating the conformation of the unfolded state in a lipid bilayer, which provides a more defined hydrophobic environment than micelles, will be a crucial future task to understand thermodynamics and mechanisms of membrane protein folding in cell membranes.

We identified subglobal unfolding of the C-terminal region which encompasses the active site. This asymmetric stability profile of GlpG is analogous to the highly-polarized folding transition state possessing a compact folding nucleus in TM1–TM2 and largely unstructured TM3–TM6\(^{24}\). Single-molecule tweezers study has also identified TM3–TM6 or TM5–TM6 as a flexible region\(^{25}\). Although we defined the region that underwent subglobal unfolding as the approximate C-terminal half, it would be more reasonable to interpret it as an ensemble-averaged event which involved unfolding of a various number of the C-terminal helices. Our work is unique in that we demonstrated partial unfolding even under a non-denaturing condition, which reflects intrinsic conformational malleability of the region that encompasses the active site. Although it is not clear if subglobal unfolding is necessary during the catalytic cycle of GlpG, we speculate that this malleability is adequate for conformational changes required for substrate interaction and catalytic mechanism. Further supporting this idea, disordering of the L5 loop, partial unfolding of TM5 and tilting of TM6 have been observed from crystal structures of GlpG in apo and inhibitor-bound forms\(^{27,46,47}\).

Our unprecedented cooperativity analysis suggests that the helical-bundle architecture of GlpG is maintained through a network of cooperative and localized interactions. Although the cooperativity network and its role in protein stability and function have been analyzed for water-soluble proteins\(^{4,7,9,41,48}\), such aspects have not been investigated for membrane proteins. Our experimentally-determined cooperativity map indicates that the degree of cooperativity was the largest for the buried residues near the center of the membrane and faded out towards the lipid- and water-contacting regions. This positional dependence of the
cooperativity profile suggests that complex environmental constraints for stabilizing membrane proteins, \textit{i.e.} protein-protein, protein-lipid and protein-water interactions, play an important role in the organization of the interaction network. Our general steric trapping strategy and steric trapping-based approaches will serve as powerful tools for exploring the folding energy landscape of membrane proteins in native lipid bilayers, which still remains as a far-reaching goal.

**ONLINE METHODS**

**Synthesis of BtnPyr-IA and BtnRG-TP**

Synthesis schemes and characterizations are shown in Supplementary Notes.

**Preparation of GlpG DNA constructs**

GlpG gene was amplified from chromosomal DNA of \textit{E. coli} strain MG1655 (Coli Genetic Stock Center at Yale University) using primers containing \textit{NdeI} and \textit{BamHI} restriction sites. The amplified gene was ligated into pET15b vector with an N-terminal His$_6$-tag. Site-directed mutagenesis for introducing amino acid substitutions was performed using the QuikChange Site-Directed Mutagenesis Kit (Agilent).

**Expression and purification of GlpG**

GlpG was expressed in \textit{E. coli} BL21(DE3) RP strain. Cells were grown at 37 °C until OD$_{600}$ = 0.6 was reached. Protein expression was induced with 0.5 mM isopropyl \textbeta-\thiogalactopyranoside (IPTG, GoldBio), followed by additional cultivation at 15 °C for 16 h. GlpG was purified from the total membrane fraction obtained by ultracentrifugation (Beckman Coulter, Type 45 Ti rotor, 50,000 g, for 2 h) using Ni$^{2+}$-NTA affinity chromatography (Qiagen) after solubilization with 2% n-dodecyl-\textbeta-D-maltoside (DDM, Anatrace).

**Labeling of GlpG and determination of labeling efficiency using SDS-PAGE gel shift assay**

For labeling, purified cysteine variants (0.2% DDM, 50 mM Tris-(hydroxymethyl) aminomethane hydrochloride (TrisHCl), 200 mM NaCl and pH 8.0) were diluted to less than 100 μM and incubated with a ten-fold molar excess Tris(2-carboxyethyl)phosphine hydrochloride (TCEP-HCl, Pierce) for 1 h at room temperature. 40 times molar excess of BtnPyr-IA or BtnRG-TP dissolved in dimethyl sulfoxide (DMSO) (~20 mg/ml) was added while vortexing. Labeling reaction was allowed to proceed at room temperature overnight in the dark. Excess free labels were removed by extensive washing of the proteins bound to Ni$^{2+}$-NTA affinity resin using 0.2% DDM, 50 mM TrisHCl, 200 mM NaCl and pH 8.0 solution. Labeled GlpG was dialyzed against 0.02% DDM, 50 mM TrisHCl, 200 mM NaCl, pH 8.0 buffer to remove imidazole.

Typically, the labeling efficiency of BtnPyr-IA and BtnRG-TP ranged from 1.5–2.2 as estimated from SDS-PAGE gel shift assay or comparison of the concentration of BtnPyr determined by pyrene absorbance ($\varepsilon_{346\text{nm}}=43,000 \text{ M}\cdot\text{cm}^{-1}$) and the concentration of GlpG determined by DC protein assay (Bio-Rad). SDS-PAGE was employed utilizing the facts that mSA maintains its tetrameric structure and the biotin-mSA complex is resistant to

*Nat Chem Biol.* Author manuscript; available in PMC 2016 September 21.
dissociation in the presence of SDS. SDS-PAGE gel shift assay (Supplementary Fig. 14) was carried out as follows: 10 μl of 5 μM of labeled GlpG was incubated with 10 μl of 2% SDS sample-loading buffer with 10% (v/v) β-mercaptoethanol for 30 min. Then, wild-type monovalent streptavidin (mSA-WT) was added to labeled GlpG (GlpG:mSA-WT molar ratio of 1:3) and the mixture was incubated at room temperature for 30 min before SDS-PAGE without sample heating. The gel box was incubated in ice during electrophoresis to prevent heat-induced dissociation of mSA-WT bound to biotin label on GlpG. Labeling efficiency was determined by comparing the intensities that correspond to single-mSA bound GlpG and double-mSA bound GlpG after accounting for the molecular mass of GlpG and mSA (AlphaImager, ProteinSimple). GlpG with no label was not considered because this species does not bind mSA, thus not participating in steric trapping. mSA was prepared as described previously.

Expression and purification of GlpG substrate SN-LacYTM2

As a folding indicator for GlpG, we used its proteolytic activity mediating specific cleavage of a transmembrane (TM) substrate, the second TM domain of the lactose permease of E. coli fused to staphylococcal nuclease (SN-LacYTM2) (Supplementary Fig. 1). The DNA construct for LacYTM2 was amplified from a DNA template containing full length lactose permease using primers containing XmaI and XhoI restriction sites, which was then ligated into a pET30a vector containing SN domain, TEV protease recognition site, and C-terminal His6-tag (SN-TEV-LacYTM2-His6). In the LacYTM2 region, the position which was five residues upstream from the scissile bond (P5 position) was substituted with cysteine for labeling with thiol-reactive, environment-sensitive fluorophore iodoacetyl-7-nitrobenz-2-oxa-1,3-diazol (IA-NBD amide, Setareh Biotech). SN-TEV-LacYTM2-His6 containing the substituted cysteine was expressed in BL21(DE3) RP E. coli strain. The protein was expressed, purified and labeled using the protocol for SN-GpATM-His6 described previously.

Fluorescence-based high-throughput activity assay for GlpG

Activity assay (Supplementary Fig. 1) was initiated by addition of 10 times molar excess of NBD-labeled SN-LacYTM2 to purified GlpG. Time-dependent changes of NBD fluorescence was monitored in 96-well plate using SpectraMax M5e plate reader (Molecular Devices) with excitation and emission wavelengths of 485 nm and 535 nm, respectively. Fluorescence change was normalized to a control sample containing NBD-SN-LacYTM2 alone. The effect of mSA binding on the activity of single- and double-biotin variants of GlpG were tested by addition of excess mSA-WT (20 μM). The GlpG-mSA mixture was incubated overnight (single-biotin variants), for 2 days (172/267C-BtnPyr2) or for 5 days (95/172N-BtnPyr2). Folding reversibility was tested using the following steps: Each double-biotin GlpG variant was first inactivated with mSA-S27A variant (20 μM) possessing a weaker biotin binding affinity (KD,bioin=1.4 nM) for the same incubation time as with mSA-WT. Next, excess free biotin (2 mM) was added to induce competitive dissociation of bound mSA. The activity of refolded GlpG was measured after incubation overnight. The statistical significance of the activity changes upon unfolding and refolding were evaluated using Student’s t-test (n=3–5).
Double electron-electron resonance EPR spectroscopy (DEER-EPR)

DEER-EPR measurements were performed on a Bruker Elexsys 580 spectrometer with Super Q-FTu Bridge, Bruker ER 5107DQ resonator and 10 W Q-band amplifier at 80 K. The spin-labeled samples ranging from 80 to 160 μM GlpG were flash-frozen in quartz capillaries using a liquid nitrogen bath immediately prior to data collection. For data collection, 36-ns π-pump pulse was applied to the low field peak of the nitroxide absorption spectrum, and the observer π/2 (16 ns) and π (32 ns) pulses were positioned 17.8 G (50 MHz) upfield, which corresponded to the nitroxide center resonance. A two-step phase cycling (+x, −x) was carried out on the first (π/2) pulse from the observer frequency. The time domain signal collected for each sample varied from 2.3 to 2.5 μs. Based on the collection time, the reliable inter-spin distance range was ~15−~60 Å. DEER data were analyzed using the program LongDistances, which was written in LabVIEW by Christian Altenbach (http://www.biochemistry.ucla.edu/biochem/Faculty/Hubbell/).

Time-dependent proteolysis of GlpG using chymotrypsin

GlpG variants (5 μM) were incubated for 7 days (95/172N-BtnPyr2: residual activity 40%) or 4 days (172/267C-BtnPyr2: residual activity 20%) in the presence and absence of 25 μM mSA-WT in 20 mM sodium phosphate (pH 7.5), 20 mM DDM, 200 mM NaCl, 1 mM TCEP buffer. In the presence of mSA-WT, the residual activities were ~40% for 95/172N-BtnPyr2 and ~20% for 95/172N-BtnPyr2 relative to those without mSA before proteolysis. Proteolysis of 5 μM casein (from bovine milk, Sigma) was performed in the same buffer condition but without prolonged incubation. For all protein samples, proteolysis was initiated by the addition of 2.5 μM chymotrypsin (bovine chymotrypsin-α: sequencing grade, Sigma) to 10 μL aliquots, and quenched at specified time by the addition of 10 mM permethylsulfoxide. Time-dependent proteolysis was monitored by SDS-PAGE (Supplementary Fig. 7).

Construction of binding isotherms to determine thermodynamic stability of GlpG by steric trapping using FRET

1 μM of GlpG labeled with BtnPyr was titrated with mSA specifically labeled with DABCYL-plus-maleimide (AnaSpec) at Y83C-position of the active subunit (mSA_{DAB}) in 5 mM DDM, 0.25 mM TCEP, 20 mM sodium phosphate and 200 mM NaCl (pH 7.5). The titrated samples were transferred to a 96-well UV-compatible microplate, sealed with a polyolefin tape, and incubated for 5 days (for 95/172N-BtnPyr2) or 2 days (for 172/267C-BtnPyr2) at room temperature. Binding was monitored by the decrease of pyrene-monomer fluorescence at 390 nm with an excitation wavelength of 345 nm using SpectraMax M5e plate reader. Data were averaged from four readings. Nonspecific FRET was obtained by measuring the fluorescence intensity of double-biotin GlpG variants which were pre-saturated with 10 μM of the high-affinity mSA-WT (without DABCYL-label) at an increasing concentration of the lower-affinity variant mSA_{DAB}-S45A. In this condition, mSA_{DAB}-S45A cannot compete for biotin label and only diffuses around in the solution.
Fitting of binding isotherm to obtain thermodynamic stability of GlpG

Fitting equation to obtain thermodynamic stability of GlpG using steric trapping was based on the following reaction scheme:

\[ F \cdot mSA \rightleftharpoons U \cdot mSA, \quad \text{where} \quad K_U = \frac{[U \cdot mSA]}{[F \cdot mSA]} \quad (2) \]

\[ U \cdot mSA + mSA \rightleftharpoons U \cdot 2mSA, \quad \text{where} \quad K_{d,\text{biotin}} = \frac{[U \cdot mSA][mSA]}{[U \cdot 2mSA]} \quad (3) \]

Fitting equation for the second mSA binding phase was:

\[ F = \frac{1}{1 + \left( \frac{K_{d,\text{biotin}}}{K_U} \right) \left( \frac{1}{[mSA]} \right)} (F_\infty - F_0) + F_0 \quad (4) \]

where \( F \) is measured fluorescence intensity, and \( F_0 \) and \( F_\infty \) are the fluorescence intensities from GlpG labeled with BtnPyr at [mSA]=0 and at the saturated bound level, respectively. [mSA] is the total mSA concentration, \( K_{d,\text{biotin}} \) is the dissociation constant for unhindered biotin binding affinity of mSA (Supplementary Fig. 3), and \( K_U \) is the equilibrium constant for unfolding of GlpG. After obtaining the fitted \( K_U \), the thermodynamic stability was calculated using the equation \( \Delta G_{\text{U,ST}} = -RT\ln K_U \).

Determination of biotin affinity (\( K_{d,\text{biotin}} \)) of mSA variants by FRET

Biotin binding affinity of a weaker binding mSA variant mSA-W79M was measured by titration of 50 nM GlpG single cysteine variants labeled with BtnPyr (FRET donor) with mSA\textsubscript{DAB}-W79M (FRET acceptor) in 5 mM DDM, 0.25 mM TCEP, 20 mM sodium phosphate and 200 mM NaCl (pH 7.5). The titrated samples were transferred to a 96-well UV-compatible microplate, sealed with a polyolefin tape, and incubated for 24 h at room temperature. Binding was monitored by the decrease of pyrene-monomer fluorescence at 390 nm with an excitation wavelength of 345 nm using SpectraMax M5e plate reader. Data were averaged from four readings. For fitting of binding data to obtain \( K_{d,\text{biotin}} \) of mSA\textsubscript{DAB}-W79M, the following equation was used:

\[ F = A_1 \times \left( \frac{P_T + [mSA] + K_{d,\text{biotin}}}{2P_T} \right) - \sqrt{\left( \frac{P_T + [mSA] + K_{d,\text{biotin}}}{2P_T} \right)^2 - 4P_T [mSA]} + A_2 \quad (5) \]

where \( F \) is measured fluorescence intensity, \( P_T \) is total GlpG concentration, [mSA] is the total mSA concentration (variable), \( K_{d,\text{biotin}} \) is the dissociation constant for biotin binding affinity of mSA\textsubscript{DAB}. A1 is the net fluorescence change, A2 is the fluorescence level without mSA\textsubscript{DAB}. Fitted values include \( K_{d,\text{biotin}} \), A1 and A2; other known values were fixed.
To determine $K_{d,\text{biotin}}$ of tight-binding mSA variants and mSA variants lacking DABCYL quencher label, a FRET-based competition assay was employed. 1 μM GlpG was pre-equilibrated with 2–5 times excess of mSA$_{\text{DAB}}$ variant for 1 h at room temperature. In this condition, pyrene fluorescence was suppressed. Weaker-affinity unlabeled mSA variant was titrated into the sample. Here either DABCYL-labeled or unlabeled mSA variant had known $K_{d,\text{biotin}}$. The titrated samples were transferred to a 96-well UV-compatible microplate, sealed with a polyolefin tape, and incubated for 24 h at room temperature. Resultant dissociation of mSA$_{\text{DAB}}$ by competition was monitored by the increase of pyrene-monomer fluorescence at 390 nm with an excitation wavelength of 345 nm using SpectraMax M5e plate reader. Data were averaged from four readings. For fitting of competition data to obtain unknown $K_{d,\text{biotin}}$, the following equation was used:

$$F = A1 \times \frac{-\left[ P_T + [mSA] + \frac{K_{\text{unlabel}}}{K_{d,\text{DAB}}} (C_T - P_T) \right] + \sqrt{\left( P_T + [mSA] + \frac{K_{\text{unlabel}}}{K_{d,\text{DAB}}} (C_T - P_T) \right)^2 + 4P_T [mSA] \frac{K_{\text{unlabel}}}{K_{d,\text{DAB}}} + A2}}{2P_T \frac{K_{\text{unlabel}}}{K_{d,\text{DAB}}}}$$

(6)

where $F$ is measured fluorescence intensity, $P_T$ is the total GlpG concentration, $C_T$ is the total mSA$_{\text{DAB}}$ concentration, $[mSA]$ is the total unlabeled mSA concentration (variable), $K_{\text{unlabel}}$ is the $K_{d,\text{biotin}}$ for mSA without a DABCYL label, $K_{d,\text{DAB}}$ is the $K_{d,\text{biotin}}$ for mSA$_{\text{DAB}}$, $A1$ is the amplitude of binding, and $A2$ is the initial fluorescence level. Fitted values include the unknown $K_{\text{unlabel}}$ or $K_{d,\text{DAB}}$, $A1$ and $A2$; all other values are fixed.

**SDS-denaturation of GlpG variants labeled with BtnPyr**

0.4 uM GlpG doubly-labeled with BtnPyr was titrated with SDS in 5 mM DDM, 20 mM Na$_2$HPO$_4$, 200 mM NaCl, pH 7.5, ranging from 0–0.9 SDS mole-fraction ($X_{\text{SDS}}=[\text{SDS}] / ([\text{SDS}]+[\text{DDM}])$. Samples were incubated overnight at room temperature. The detailed scheme for fluorescence spectroscopy to monitor SDS-induced equilibrium unfolding is described in **Supplementary Fig. 8**. The unfolding curves were constructed using the average of three measurements. To determine thermodynamic stability of GlpG from SDS denaturation ($\Delta G_{U,\text{SDS}}^\circ$), the unfolding curves were fitted to the following two-state Santoro-Bolen equation:

$$F = \left[ (F_{\text{U}} - \frac{m_{\text{F}} X_{\text{SDS}}}}{1 + \exp \left( \frac{m_{\text{F}} X_{\text{SDS}}}}{RT} \right) } \right]$$

(7)
\[ \Delta G^0_{U,SDS} = X_{SDS/2} \times m_{SDS} \]  

(8)

\( F \) is the net fluorescence change. Baselines for the pre- and post-transition regions were determined by the fitted parameters: \( F_F \), the fluorescence value for fully folded GlpG; \( F_U \), the fluorescence value for fully unfolded GlpG; \( m_F \), the slope of the fully folded baseline; and \( m_U \), the slope of the fully unfolded baseline. \( m_{SDS} \) is the slope of the transition region against \( X_{SDS} \) and \( X_{SDS/2} \) is the transition midpoint. Fitted values include \( F_F \), \( F_U \), \( m_F \), \( m_U \), \( m_{SDS} \), and \( X_{SDS/2} \). \( \Delta G^0_{U,SDS} \) represents thermodynamic stability in the absence of denaturant.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**ACKNOWLEDGMENTS**

This work was supported by start-up fund from Michigan State University (H.H.), Hunt for a Cure grant for basic cystic fibrosis research (H.H), National Cancer Institute R01CA149541 (X.H), National Eye Institute R01EY05216 (W.L.H.) and the Jules Stein Professor endowment (W.L.H.). The authors thank John McCracken for general advice on EPR and helping with CW-EPR measurements and Christian Altenbach for DEER analysis program. We thank Erin Deans and Caleb Arthur for helping with protein preparations. We also thank Robert Jefferson, Nicholas Woodall, Yu-Chu Chang, David Weliky, Lee Kroos, John McCracken and James Bowie for critical reading of the manuscript. We appreciate suggestions from anonymous reviewers, which greatly helped improving the quality of this manuscript.

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Figure 1. Principle of steric trapping and steric-trapping probes developed in this study
(a) Steric trapping principle for measuring thermodynamic stability ($\Delta G^\circ_{\text{U}}$) of proteins. After conjugation of biotin tags to two specific residues that are spatially close in the folded state but distant in the amino acid sequence, the first mSA binds unhindered to either biotin label with intrinsic binding affinity ($\Delta G^\circ_{\text{Bind}}$). Due to the steric hindrance with pre-bound mSA, the second mSA binds only when the native tertiary contacts between biotinylated sites are unraveled by transient unfolding. Coupling of mSA binding to unfolding leads to attenuation of the apparent binding affinity of the second mSA relative to that of the first.
mSA, whose degree is correlated with the protein stability. Thus, thermodynamic stability of the target protein can be determined by fitting of the second binding phase (see equations (2)–(4) in Online Methods). Overall, protein unfolding is driven by the affinity and concentration of mSA without perturbing the native solvent condition. Folding reversibility is tested upon addition of excess free biotin by which bound mSA molecules are released by competition. (b) Thiol-reactive biotin derivatives possessing a spectroscopic reporter group developed in this study. BtnPyr-IA (1): biotin (red shaded)-pyrene (green shaded)-iodoacetamide (blue shaded) conjugated to a lysine template, and BtnRG-TP (2): biotin (red shaded)-1-oxyl-2,2,5,5-tetramethylpyrroline spin label (green shaded)-thiopyridine (blue shaded) conjugated to a cysteine template.
Figure 2. GlpG reversibly unfolds by double-binding of mSA

(a) Locations of two different biotin pairs for steric trapping in the structure of GlpG (PDB code: 3B4549) and their C\textalpha-C\textalpha distances. (b) Reversible control of GlpG folding tested by the proteolytic activity as a folding indicator. The second TM domain of the lactose permease50 of E. coli fused to staphylococcal nuclease was used as a model substrate throughout this study (Supplementary Fig. 1). All activity levels were normalized relative to the activity of wild-type GlpG. Error bars denote mean ± s. d. (n=5 for the data without mSA and n=3 for the data with mSA). **Top panels**: binding of wild-type mSA (mSA-WT) to
individual single-cysteine variant labeled with BtnPyr did not affect the activity. **Bottom panels:** saturated binding of mSA-WT to each double-cysteine variant labeled with BtnPyr led to an inactivation of GlpG (the second bar from the left in each panel). To test folding reversibility, double-biotin GlpG variants were first inactivated with mSA-S27A possessing a weaker biotin binding affinity ($K_{d,\text{biotin}}=1.4\times10^{-9}$ M) for 2–5 days (the third bar). Next, excess free biotin was added to induce competitive dissociation of bound mSA (the fourth bar). All $p$-values obtained from Student’s $t$-test were smaller than the threshold significance level ($\alpha=0.05$), indicating that the activity changes for the unfolding and refolding reactions were significant.
Figure 3. DEER suggests steric trapping induce wide separation of two biotinylated sites

Background-subtracted dipolar evolution data and their fits (left) and inter-spin distances (right) for the native (dashed lines), SDS-induced unfolded (gray solid lines, SDS mole fraction = \([\text{SDS}] / ([\text{DDM}] + [\text{SDS}]) > 0.8\), in which the unfolded fraction exceeded 0.9), and steric-trapped (black solid lines) unfolded states for (a) 95/172N-BtnRG2 GlpG and (b) 172/267C-BtnRG2 GlpG. The approximate upper limit of the reliable mean distance was \(\sim 53 \text{ Å}^{32}\).
Figure 4. Thermodynamic stability of GlpG using steric trapping and SDS denaturation
(a) Steric-trapping strategy using FRET between fluorescent pyrene (donor) in BtnPyr labeled on GlpG and non-fluorescent quencher DABCYL (acceptor) thiol-specifically labeled near the biotin binding pocket (Y83C) of the active subunit of mSA (mSA_{DAB}). (b) Equilibrium unfolding of GlpG variants 95/172_{N}-BtnPyr$_2$ and 172/267$_{C}$-BtnPyr$_2$ as a function of SDS mole fraction measured by FRET between Trp residues (donor) of GlpG and pyrene (acceptor) on BtnPyr labels (Supplementary Fig. 8). Errors in $\Delta G^0_{U,SDS}$ values denote mean ± s. d. from fitting. (c) Binding isotherms of 95/172$_{N}$-BtnPyr$_2$ and 172/267$_{C}$-
BtnPyr2 with three mSA variants mSA_{DAB-WT} (black circles, $K_{d,\text{biotin}}=\sim 10^{-14}$ M), mSA_{DAB-S27A} (red circles, $K_{d,\text{biotin}}=1.4\times10^{-9}$ M) and mSA_{DAB-S45A} (blue circles, $K_{d,\text{biotin}}=9.0\times10^{-9}$ M) (Supplementary Fig. 3). The activity change for each double- biotin variant (crosses, right y-axis) was measured at an increasing concentration of mSA_{DAB-S27A}. The thermodynamic stability ($\Delta G_{U,ST}$) of each variant was obtained by fitting of the second mSA-binding phase to equation (4) in Online Methods. Procedures to obtain nonspecific FRET (open circles) are described in Online Methods. Errors in fluorescence denote mean ± s. d. (n=4). Errors in activity denote ± s. d. from fitting. Errors in $\Delta G_{U,ST}$ values denote mean ± s. d. (n=3).
Figure 5. Dependence of thermodynamic stability of GlpG on SDS mole fraction
The plot containing $\Delta G^o_{U,ST}$'s (diamonds) obtained by steric trapping and $\Delta G^o_{U,SDS}$'s (squares) obtained by SDS denaturation as a function of SDS mole fraction ($X_{SDS}$) for 95/172N-BtnPyr$_2$ and 172/267C-BtnPyr$_2$. To fit $\Delta G^o_{U,ST}$, we accounted for the changes in the biotin affinity of mSA$_{DAB}$ variants which depended on $X_{SDS}$ (Supplementary Fig. 3d). Errors in $\Delta G^o_{U,ST}$ denote ± s. d. from fitting. Solid lines are the linear-regression fits of $\Delta G^o_{U,ST}$ in the range of $X_{SDS}=0.2-0.4$ and dashed lines indicate the extrapolation lines of $\Delta G^o_{U,SDS}$ to zero $X_{SDS}$ from equilibrium SDS denaturation. The slope in the $\Delta G^o_{U}$ vs $X_{SDS}$ plot represents the $m$-value. For 95/172N-BtnPyr$_2$, $m=16±3$ (blue dashed line) from SDS denaturation and $m=14±2$ (blue solid line) from steric trapping. For 172/267C-BtnPyr$_2$, $m=17±2$ (red dashed line) from SDS denaturation and $m=8±1$ (red solid line) from steric trapping. Errors in the $m$-values denote ± s. d. from fitting.
Figure 6. Cooperativity map reveals a network of clustered cooperative and localized interactions for the stability of GlpG under a native micellar condition
(a) Scheme for quantifying the cooperativity of interactions of a specific side chain. The stability changes ($\Delta\Delta G^\circ_U$) induced by the same mutation (black star) were probed with two biotin pairs, 95/172$^N$-BtnPyr$_2$ and 172/267$^C$-BtnPyr$_2$ located in the N- and C-terminal regions, respectively, and compared to each other to yield $\Delta\Delta\Delta G^\circ_U$ using equation (1). The cyan-backbone region designates subdomain I (TM1-L1-TM2-TM3-L3$_{198}$), which ends at residue 198 in the L3 loop (marked with a magenta wedge) and the yellow-backbone region
(L319-TM4-TM5-L5-TM6) indicates subdomain II. The uncertainty of the subdomain-division point was ±20–30 residues around residue 198. The detailed strategy for the subdomain dissection and the estimation of the uncertainty is described in Supplementary Fig. 10. Catalytic dyad composed of Ser201/His254 is shown as spheres. (b) Cooperativity map at a side-chain resolution. The map shows the “cooperative” (green, $|\Delta\Delta\Delta G_U|<RT=0.6 \text{ kcal/mol}$) and “localized” ($|\Delta\Delta\Delta G_U|>RT$) side-chain interactions. Localized interactions were further divided using additional cut-off energy values, $2RT<|\Delta\Delta\Delta G_U|<RT$ (“moderately-localized” interactions) and $|\Delta\Delta\Delta G_U|>2RT$ (“highly-localized” interactions). Each side chain was color-coded based on these criteria for $\Delta\Delta\Delta G_U$ as shown in the figure. Interactions mediated by residues G261 and A265 (denoted with stars) were “over-propagated”. Errors in individual $\Delta G_U$ were ±0.1–±0.2 kcal/mol (mean ± s. d. from fitting) and errors in $\Delta\Delta\Delta G_U$ ranged from ±0.1–±0.4 kcal/mol, which were calculated using the propagation of errors in $\Delta G_U$ (Supplementary Table 1).