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Epistasis on the stability landscape of de novo TIM barrels explored by a modular design approach

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Main Text.
Figures 1 to 5.
Table 1.
Abstract

The design of stable proteins with custom-made functions is a major goal in biochemistry with practical relevance for our environment and society. High conformational stability lowers protein sensitivity to mutations and changes in the environment; thus, understanding and manipulating protein stability will expand the applications of de novo proteins. Since the \((\beta/\alpha)_{\beta}\)-barrel or TIM-barrel fold is one of the most common functional scaffolds, in this work we designed a collection of stable de novo TIM barrels (NovoTIMs), using a computational fixed-backbone and modular approach based on improved hydrophobic packing of sTIM11, the first validated de novo TIM barrel. NovoTIMs navigate a region of the stability landscape previously uncharted by natural TIM barrels, with variations spanning 60 degrees in melting temperature and 25 kcal per mol in conformational stability throughout the designs. Significant non-additive or epistatic effects were observed when stabilizing mutations from different regions of the barrel were combined. The molecular basis of epistasis in NovoTIMs appears to be related to the extension of the hydrophobic cores. This study is an important step towards the fine-tuned modulation of protein stability by design.

Significance Statement

\textit{De novo} protein design expands our knowledge about protein structure and stability. The TIM barrel is a highly relevant fold used in nature to host a rich variety of catalytic functions. Here, we follow a modular approach to design and characterize a collection of \textit{de novo} TIM barrels and subjected them to a thorough folding analysis. Non-additive effects modulate the increase in stability when different regions of the barrel are mutated, showing a wide variety of thermodynamic properties that allow them to navigate an unexplored region of the stability landscape found in natural TIM barrels. The design of stable proteins increases the applications of \textit{de novo} proteins and provides more information on the molecular determinants that modulate structure and stability.

Main Text

Introduction

Proteins are essential macromolecules capable of performing diverse and exquisite biological functions such as nutrient uptake, environmental stimuli sensing, immune protection, energy storage, cellular communication, molecule transportation, or enzymatic reactions. To guarantee such activities, the functional states must act under specific environmental conditions in a relevant time scale, that is, proteins must be “stable”. Protein stability is required to maintain functional structures and it enhances the ability of proteins to evolve new properties \((1, 2)\). The central role of proteins in the chemistry of life, as well as their increasing application in basic and applied research, implies that the understanding and manipulation of protein stability are both practically and academically relevant.

There are two main indicators of protein conformational stability at equilibrium. One is the difference of free energy between the native and unfolded states at a given temperature \((\Delta G)\), which is often obtained by chemical unfolding experiments carried out at \(25 \, ^{\circ}\text{C}\). In addition, stability is also assessed in the context of thermal unfolding, where the unfolding temperature \((T_m)\), the temperature at the midpoint of the transition from native to the unfolded state, is the most common parameter employed to quantify stability. Both the \(\Delta G\) and \(T_m\) parameters, usually determined as criteria for a “stable” protein, are related with the enthalpy \((\Delta H)\) and heat capacity
Changes through the Gibbs-Helmholtz equation, which describes the variation of $\Delta G$ with temperature, the so-called “stability curve” of proteins (3). Different mechanisms have been proposed to modify the stability curve of proteins (4), and numerous studies on natural proteins and their site-directed mutants have been used to rationalize the stability of thermophilic proteins and also to engineer thermostability (5). Historically, the design of stable proteins has been one of the main objectives of computational protein design (6). Several strategies, such as increasing the hydrophobic area in internal cores, improvement of water-protein interactions, the introduction of disulfide bridges as well as the addition of salt bridges, have been proposed (7-18). The design of de novo proteins can further enhance our understanding of the physicochemical properties that modulate stability. For example, although folding behavior has been only addressed for very few cases, the kinetic analysis of the folding mechanism of two de novo $\beta\alpha$ proteins has revealed complex free energy surfaces (19, 20). The fine-tuning of conformational stability, that is, the manipulation of the protein stability curve, is an open challenge for protein design and engineering. Such a goal requires a comprehensive characterization of de novo proteins, describing the combination of thermodynamic parameters that can be reached in a particular fold.

Within the different topologies that a protein can adopt, the TIM-barrel or ($\beta/\alpha$)$_8$-barrel fold is one of the most abundant superfolds in nature, since $\sim$10% of all known proteins adopt this structure (21). Furthermore, proteomic analysis shows that the TIM-barrel domain is within the mean size of the proteins present in Escherichia coli (22). Besides, the TIM-barrel fold is one of the most successful topologies used in nature to host catalytic activities. Due to its large variety of functions and its ubiquity in different types of enzymes, the TIM barrel represents a very suitable scaffold for protein function design and engineering, with numerous applications in the fields of synthetic biology, diagnosis, and imaging (23). For these reasons, its construction has been an important objective for protein design over the years (24–28). Recently, the successful design of a de novo four-fold symmetric TIM barrel was described: the sTIM11 protein (29). Considering that sTIM11 presents a sequence distant from the naturally occurring TIM-barrel superfamilies, the potential of the TIM-barrel fold to bear functions is more significant than we know so far. sTIM11 shows a high melting temperature ($T_m$ = 80 °C) but low conformational stability ($\Delta G_{25^\circ C} = \sim$4 kcal mol$^{-1}$) when compared to natural TIM barrels (30-32). Since low conformational stability often results in high sensitivity to mutations and changes in the environment, this can limit the design of novel proteins with new functions (8). Thus, fine-tuning the stability of the sTIM11 scaffold is a prerequisite to functionalize and generate tailor-made barrels for applications in biochemistry, biotechnology, and medicine. In this work, a fixed-backbone design with a modular approach was used to generate a collection of de novo TIM barrels. Their stability landscape and structural properties were characterized in detail increasing our knowledge on how stability can be fine-tuned by design.

Results and Discussion

Modular repacking of the TIM-barrel hydrophobic cores

The de novo protein sTIM11 is an idealized four-fold symmetric TIM barrel of 184 residues, which was designed to include two cysteines that, however, did not form the intended disulfide bond (Fig. 1). To avoid reactive free thiols, both residues were reverted to the residues in the original four-fold design (C8Q and C181V), resulting in sTIM11noCys. The base design NovoTIM0, which is the starting point for all further constructs in this work, additionally contains the changes W34V and A38G in all symmetry-related quarters. These residues are situated in every second $\alpha/\beta$-loop, and in sTIM11, these tryptophan residues are the most highly solvent exposed. While different strategies have been explored to increase protein stability (8, 18), here we focused on hydrophobic repacking. The structural analysis suggested two regions to be amenable to improvements in sTIM11, namely the central and the peripheral hydrophobic cores. The interior of the circular sheet forms the central core, whereas the outer face of the strands and the internal face of the helices constitute the peripheral core. In this latter, we identified two regions with
internal cavities that are located in the lower and upper parts of the barrel, respectively (Fig. 1). The residues lining the three aforementioned regions were subjected to fixed-backbone Rosetta design according to the flow diagram depicted in Fig. S1.

Ten designs were selected for characterization in the first round: four designs with modifications in the internal core (NovoTIM1-4) as well as three designs each for the bottom peripheral or outer core (NovoTIM5-7) and the top peripheral core (NovoTIM8-10) (Fig. S2). For the inner core, it was not possible to find better designs when four-fold symmetry was preserved. Therefore, in NovoTIM1-4, as well as all the designs that contain them, only a two-fold symmetry was enforced. An exploratory characterization by circular dichroism (CD) and differential scanning calorimetry (DSC) of the proteins from the first design round (NovoTIMs 1-10) as well as NovoTIM0 showed that NovoTIM1, NovoTIM6, and NovoTIM8 were the best designs of each region (Fig. S3 and supplementary text S1).

To test for additivity effects on stability and structure, mutations contained in the best design of each group were combined to generate the following double-region designs: NovoTIM11 (NovoTIM1 + NovoTIM6), NovoTIM12 (NovoTIM1 + NovoTIM8), and NovoTIM13 (NovoTIM6 + NovoTIM8). Finally, in the third design round the mutations of all three regions were combined resulting in NovoTIM14 (NovoTIM1 + NovoTIM6 + NovoTIM8). All these proteins as well as sTIM11, sTIM11noCys, and NovoTIM0 were characterized in detail. Sequences, mutations in each design, and sequence identity data are reported in supplementary information (Fig. S2 and tables S1-S4).

**Physicochemical characterization of NovoTIMs**

sTIM11, sTIM11noCys, and all NovoTIM variants presented the characteristic far-UV CD spectra observed for α/β proteins (Fig. 2A and Fig. S4). Accordingly, the near-UV CD and intrinsic fluorescence (IF) spectra showed that the aromatic residues are buried from the solvent and structured in the folded state (Fig. 2B-2C and Fig. S5-S6; see supplementary text S2 for details). All NovoTIMs adopt a monomeric and compact shape as revealed by the invariant value of the Stokes radius determined by analytical size exclusion chromatography over a twenty-fold protein concentration range (table S5). NovoTIM0 is also monomeric but shows a Stokes radius (26.1 ± 0.3 Å) slightly higher than that expected for a compact protein of this size, but still far away from the expected value for an unfolded conformation (22.5 ± 1.0 Å and 42.0 ± 1.0 Å, respectively; 33). This is in agreement with the red shift in the IF spectra and suggests a slightly expanded conformation for NovoTIM0.

Thermal unfolding was then studied by CD and DSC (Fig. 2D-2E). All NovoTIMs showed cooperative transitions with a remarkably broad range of $T_m$ values, from 47 °C (NovoTIM0) to 109 °C (NovoTIM12) (table 1); indeed at 90 °C many of the proteins still showed secondary and tertiary structure (Fig. S4B and Fig. S5B). All NovoTIMs endotherms, except NovoTIM13 and NovoTIM14, showed reversibility and were well fitted to the two-state model (N $\leftrightarrow$ U) (Fig. S7 and table 1). This is remarkable because the temperature-induced unfolding of natural proteins of this size, particularly TIM barrels, is usually not reversible (22, 30). NovoTIM14 showed two endotherms, suggesting the presence of an unfolding intermediate (Fig. S7I). For NovoTIM13 and NovoTIM14, endotherms were well-fitted to an irreversible two-state mechanism (N $\rightarrow$ F) giving activation energies of 118 and 37.5 kcal mol$^{-1}$, respectively, resulting in very different kinetic stabilities (Fig. S8-S9 and supplementary text S3).

The observed unfolding $\Delta H$ and $\Delta C_p$ also vary greatly (table 1); for some NovoTIMs these values are similar to the ones expected for a protein of 184 residues, whereas for others they are smaller ($\Delta H = 128.4 \pm 3.5$ kcal mol$^{-1}$ and $\Delta C_p = 2.6 \pm 0.04$ kcal mol$^{-1}$ K$^{-1}$, according to parametric equations reported in 34). The $\Delta H$ values observed for the first and second design rounds (0.24 to 0.64 kcal mol$^{-1}$ residue$^{-1}$) are similar to those reported for natural monomeric TIM barrels (0.25 to 0.67 kcal mol$^{-1}$ residue$^{-1}$). A decrease in $\Delta C_p$ has been shown to result from residual structure in the unfolded state (35). This is observed in the far-UV CD spectra of those NovoTIMs that are
All the selected first- and second-round designs presented a ∆G at 25 °C higher than NovoTIM0, whereas the triple-design, NovoTIM14, showed a pronounced increase in stability (∆GIC = 26.7 kcal mol⁻¹; table 1). For NovoTIM14, the stability change related to the loss of the native state (∆GNi = 9.3 kcal mol⁻¹) is similar to the ∆G of the second-round designs, whereas the stability of the intermediate is much higher (∆GI-U = 17.4 kcal mol⁻¹). For three-state folders, the change in free energy from the native to the intermediate state (∆GI-I) has been termed “the relevant stability” because the intermediate is expected to be non-functional, whereas the stability change from the intermediate to the unfolded state (∆I-U) is designated as “residual stability” (36).

The modular design approach used in this work improved both ∆G and Tm substantially and hence produced significantly more stable proteins, particularly in the second- and third-round designs. In this context, it is worth mentioning that over the years the combination of stabilizing mutations has been considered an effective strategy to enhance the stability of small proteins (36, 40-43). Previous work on small globular proteins with optimized hydrophobic cores and interactions on the surface exhibited increased thermal stability by up to 30 degrees (9, 11, 15). Extending these strategies from point mutants to regions appears to be useful for bigger folds such as the TIM barrel. In what follows, using the thermal and chemical unfolding data described above, the thermodynamic properties underlying the stability of NovoTIMs are analyzed.

Global thermodynamic stability and non-additive effects of NovoTIMs

As observed in natural proteins, the m values obtained from the chemical unfolding of sTIM11, sTIM11noCys, NovoTIM0, NovoTIM6, and NovoTIM8 correlate with their ∆CP values determined by thermal unfolding (Fig. 3A), likely because both depend on the ∆ASA upon unfolding. In contrast, ∆CP values obtained for NovoTIM1, NovoTIM11, and NovoTIM12 are much lower than those expected from the reported correlation between m values and ∆CP (Fig. 3A). According to
the Rosetta models and the native state structures (see below), these differences are not exclusively due to properties of the native state since the calculated ∆ASA is close to the expected value for the size of NovoTIMs (17 135 Å²; 37). This suggests that the unfolded state reached at high temperatures is more structured than the one obtained by chemical unfolding.

The fact that many NovoTIMs show reversible temperature-induced unfolding allowed the assessment of their stability curves using the thermodynamic parameters obtained by DSC data (Fig. 3B). The ∆G values at 25 °C are in excellent agreement with those obtained from chemical unfolding experiments. According to the Gibbs-Helmholtz equation, conformational stability is modulated by changes in T_m, ∆H, and ∆C_p. For natural TIM barrels, it has been observed that changes in the stability curve are influenced mainly by modifying one or two of those parameters (30, 31). In contrast, the NovoTIMs differ in all three parameters. Increasing ∆H is the most commonly found mechanism for stabilization of thermophilic proteins (5) and is also the most often exploited mechanism for engineering protein stability (7, 40). In NovoTIMs, this mechanism is used in all proteins but is especially important in NovoTIM6, that has the highest ∆H and, therefore, a ∆G higher than NovoTIM0, indicating an enthalpy-driven stabilization (Fig. 3B). Nevertheless, in the absence of a high-resolution structure (see below), it is difficult to rationalize how enthalpic stabilization was achieved in NovoTIM6 because considerable structural rearrangements take place when new interactions are introduced or molecular strain is removed. ∆C_p determines the magnitude of the curvature of the stability curve so that changes in this parameter triggers a more or less flattened curve. A decrease in ∆C_p has been postulated as a mechanism for thermostabilization (35, 44). For NovoTIMs, the reduction in ∆C_p combined with an increase in ∆H is the reason for the increase in both T_m and stability at 25 °C. The results presented here indicate that, as observed for natural proteins, in addition to the native state, the unfolded ensemble plays an important role in shaping the stability curve and should be considered in protein design.

NovoTIMs show a non linear correlation between ∆G at 25 °C and T_m, the most commonly used parameters that describe protein stability (inset in Fig. 3C). A similar trend between ∆G at the temperature where it is a maximum (∆G_Tmax) and T_m has also been reported for natural and engineered proteins with different sizes and topologies (34, 45, 46). Additionally, the global thermodynamic stability can be conveniently described by the area (from 0 °C to T_m) under the stability curve (A). The advantage of A over ∆G at a given temperature is that A integrates the conformational stability in a temperature range (47). The relative global stability of NovoTIMs (A/ANovoTIM0) is also correlated with T_m (Fig. 3C). Notably, for NovoTIM6, NovoTIM11, and NovoTIM12, A/ANovoTIM0 is nearly ten-fold higher than for NovoTIM0 (Fig. 3C and table 1). The modular strategy used to generate the NovoTIMs and the determination of their stabilities allowed us to calculate the contribution of each region to global stability, and also to evaluate the presence of non-additive effects between different regions of the barrel. Non-additive effects were evaluated as ∆∆G_int through an approach based on thermodynamic double mutant cycles (see Materials and Methods). ∆∆G_int is also referred as to coupling energy, non-additive effect, interaction energy, and more recently epistatic effect (48). Thermodynamic cycles were constructed using the experimentally determined ∆G_{25°C} values and linking single-region/double-region designs, and then double-region/triple-region designs as indicated in Fig. S12.

It was found that stabilization is non-additive, consequently, the different barrel regions are coupled, indicating that their contribution to protein stability depends on the structural context. A positive ∆∆G_int indicates that the introduction of favorable interactions has a higher stabilizing effect when a nearby region is already mutated. All the ∆∆G_int values calculated in Fig. S12 are summarized in the single cube shown in Fig. 3D. ∆∆G_int for single and double designs (upper face of the cube) are much smaller than those observed between double- and triple-region designs. The regions that are most energetically coupled in double-region designs are the inner core (NovoTIM1) and the upper peripheral core (NovoTIM8) (∆∆G_int= 6.1 kcal mol⁻¹, see the upper panel in Fig. S12 and arrows from top to bottom face of the cube in Fig. 3D). Coupling increases considerably when a third region is incorporated on the background of two already mutated regions (∆∆G_int> 9 kcal mol⁻¹, see lower panel in Fig. S12 and arrows on the bottom face of Fig. 3D). The largest ∆∆G_int was observed when the NovoTIM8 mutations were added to NovoTIM11.
(ΔΔG_int = 17.3 kcal mol⁻¹, see purple arrow in the bottom face of Fig. 3D). Clearly, mutations in one region of the barrel can cause in a non-additive manner the loss or gain of one or more interactions in another distant region of the barrel. The latter indicates that the TIM-barrel fold is suitable for studying modularity and, in general, cooperative effects of proteins. Also, the results presented here suggest that the modular design strategy could be used in the future for the rational stability improvement in other protein topologies.

**Structural analysis of NovoTIMs**

The structural properties of NovoTIMs were examined by X-ray crystallography (table S6). High-resolution data were collected for sTIM11noCys and NovoTIM13 (1.88 and 1.64 Å, respectively), whereas a low-resolution structure was obtained for NovoTIM6 (2.90 Å). All of them showed the designed globular compact TIM-barrel topology (Fig. 4). Structural comparison of the X-ray structures and Rosetta models for sTIM11, sTIM11noCys, NovoTIM6, and NovoTIM13 showed the lowest RMSD located in the second quarter of the barrel (ranging from 0.27 to 0.68 Å). As previously observed in sTIM11 (29), the main structural differences are found in the α-helices located at the amino- and carboxyl-terminal ends. In agreement, for all the barrel structures, the RMSD among quarters of the barrel is higher in the first and fourth ones (plot in Fig. 4A). Since the TIM barrel is a closed-repeat protein, contacts between the first and last helices depend on the precise curvature generated by each α/β unit, therefore geometrical strain may interfere with the proper closure of the barrel.

A comparison of the sTIM11noCys and sTIM11 structures showed that removal of the two cysteines causes some structural changes mainly localized in the first and last quarters; the most significant deviations are observed at the amino-terminal region where the first two helices are not well-formed. So even without forming the disulfide bridge, both cysteines in sTIM11 increase the stability and promote a proper closure of the barrel (Fig. 4B and table 1). The other parts of sTIM11noCys adopt almost the same structural arrangement as in sTIM11, except for the β_6/α_7 loop which was not modeled due to an absence of electron density in that region. Thus, although removing the cysteines has effects on stability and structure, sTIM11noCys maintains the general architecture corresponding to the expected TIM barrel.

The thermodynamic properties of NovoTIM6 are very similar to those expected for a natural protein (table 1). Unfortunately, due to the low quality of the crystals and therefore the low resolution obtained (2.90 Å), details such as side-chain conformations are not well resolved in the NovoTIM6 structure. Nevertheless, it could be verified that the protein is well folded into a compact TIM-barrel (Fig. 4C). As aforementioned for sTIM11 and sTIM11noCys, when the similarity between the structure and the Rosetta model is analyzed, the four quarters in NovoTIM6 show different RMSD values (Fig. 4C). The most similar quarter is located in the second region of NovoTIM6, whereas the main deviations are located in the first and last quarter of the barrel. Almost all α/β loops of the barrel are well defined and correspond to the model. However, for some residues within 5 of the 7 β/α loops no electron density was observed. The main differences observed in the structural analysis between the Rosetta model and the NovoTIM6 structure (table S7) are likely due to the low resolution of the data where some residues and side chains are missing in the electron density map. In general, the NovoTIM6 structure has high B factors which may reflect higher disorder in the protein crystal or increased flexibility, similar to observations in some regions of sTIM11, namely the amino- and carboxyl-terminal α-helices. This could also explain difficulties in obtaining crystals that diffract at higher resolution despite many efforts (see the Materials and Methods).

As observed in all NovoTIMs, the similarities between the NovoTIM13 structure and the Rosetta model vary among the four quarters of the barrel (Fig. 4A). The second, third, and fourth quarters display minor differences between the structure and the Rosetta model, with the secondary structure elements and side chains superposing very well. The highest deviations are located at the amino-terminal region that closes the barrel (Fig. 4D). For NovoTIM13, the resolution of the
crystal structure (1.64 Å) allowed a more in-depth analysis. Most of the hydrogen bonds and salt bridges designed are observed in the NovoTIM13 structure. As a consequence of the design strategy, this number is lower than the ones for sTIM11 and stTIM11noCys, stabilizing polar interactions being replaced by an increase in hydrophobic interactions in the NovoTIM series. For example, in going from sTIM11 to NovoTIM13, a 60 % increase in the total area in hydrophobic clusters was found (3765 vs. 6148 Å²); most of this change comes from a three-fold increase in the area of the major hydrophobic cluster (1116 vs. 4351 Å²). For NovoTIM13, both the area in the major hydrophobic cluster and the total hydrophobic area found in the structure are very similar to those designed (96 and 98 %, respectively; table S7).

One of the main proposed mechanisms for the stabilization of thermophilic proteins is an increase in the number of stabilizing interactions such as salt bridges and hydrogen-bond networks (5). In fact, in going from sTIM11 to NovoTIM0, a decrease in the number of electrostatic interactions is accompanied by a decrease in stability. In contrast, the structural analysis of NovoTIMs showed that these interactions are not clearly related to the observed changes in stability. For example, some of the designs that contained the highest number of polar stabilizing interactions (such as NovoTIM1 and NovoTIM8) were not the most stable ones, whereas some of the most stable designs (such as NovoTIM6 and NovoTIMs 12-14) showed a reduction in this type of interaction (table S7). On the contrary, the stability of NovoTIMs increases with the number of hydrophobic interactions. The total area, as well as the number of residues and contacts in hydrophobic clusters, are substantially increased in the best first-round designs along with the more stable second- and third-round designs (Fig. S13 and table S7). As discussed in more detail below, this suggests that repacking of the hydrophobic cores is one of the main mechanisms to increase the thermodynamic stability of NovoTIMs.

**Epistasis on the stability landscape of de novo TIM barrels**

To correlate the most common and informative parameters obtained from both temperature and chemical unfolding, \(T_m, \Delta H,\) and \(\Delta G_{25\,\text{°C}}\) were mapped onto a “stability landscape”, a spatial representation of the observed combinations of these thermodynamic data (Fig. 5 and Materials and Methods). The \(T_m\) range found in NovoTIMs is widely distributed, covering from 47 °C to 109 °C, a range of more than 60 °C in thermostability. Besides, it was possible to design TIM barrels with stabilizing mutations that led to huge differences in stability, even higher than other systems previously reported. The stability landscape of NovoTIMs can be compared to that constructed for natural proteins (Fig. S14). The latter is rough, with some regions more populated than others, and explores an ample space due to the diversity in size, topology, oligomeric state, function, and evolutionary history of the variety of natural proteins so far characterized. Interestingly, the comparison shows that several NovoTIMs are located in a region of the stability landscape corresponding to low \(\Delta H\) and high \(\Delta G_{25\,\text{°C}}\) values, which is not populated, as far as we know, by natural proteins. The modular strategy followed in the NovoTIM design rounds can be mapped in this stability landscape. sTIM11, stTIM11noCys, NovoTIM0, and most of the first-round designs cover a vast region of the landscape valley, whereas second-round designs are located in a higher stability region. Finally, the third-round design climbs to the highest region of the landscape (Fig. 5).

Assuming additivity, the expected change in stability calculated for NovoTIM14 would be the sum of the individual stabilizations provided by all the single-region designs (NovoTIM1 + NovoTIM6 + NovoTIM8) giving a value of 9.7 kcal mol\(^{-1}\). However, the stability of NovoTIM14 is 26.7 kcal mol\(^{-1}\), indicating that more than half of the stabilization comes from positive non-additive effects. The thermodynamic cube presented in Fig. 3D shows that the \(\Delta \Delta G_{\text{int}}\) mentioned above increases in going from the first- to the second- and third-round designs. Non-additive effects or interaction energies may be referred to as epistasis, a concept traditionally used in genetics to describe the phenotype dependency of a mutation on the genetic state at other sites (48-50). Previous studies have explored and analyzed the mechanisms of epistasis within proteins, especially regarding their implications for protein function, evolution, and stability (51-55).
Rearrangements in the TIM barrel can influence local changes in other parts of the protein, and these epistatic effects are quantified in the \( \Delta \Delta G^{\text{int}} \) values whose magnitude for NovoTIMs is considerable. The structural analyses suggest that one of the molecular basis of the epistatic effect observed in NovoTIMs is likely related to the extension of the hydrophobic cores, particularly to the increase of the major hydrophobic cluster located in the interface between the inner \( \beta \)-barrel and the outer \( \alpha \)-helices (Fig. 13 and table S7). From the first- to the second-round designs, the highest area in hydrophobic clusters was found for NovoTIM12, and this corresponds to the highest positive epistatic effect in this round (\( \Delta \Delta G^{\text{int}} = 6.1 \text{ kcal mol}^{-1} \)), whereas the decrease of the hydrophobic cluster area in NovoTIM11 (compared to NovoTIM1 and NovoTIM6) correlates with a negative \( \Delta \Delta G^{\text{int}} = -1.8 \text{ kcal mol}^{-1} \). From the second- to the third-round designs, the most notable change in hydrophobic area is observed in going from NovoTIM11 to NovoTIM14, resulting in the highest positive epistatic effect (\( \Delta \Delta G^{\text{int}} = 17.3 \text{ kcal mol}^{-1} \)).

The relevance and magnitude of the epistatic or non-additive effects found in NovoTIMs, as well as those observed in other reports, suggest that modeling such interactions can improve the success in protein design and engineering.

Conclusions

Design requires a deep understanding of the relationship between sequence, structure, and stability, and therefore, the combination of thermodynamic and structural data is fundamental to achieve this goal. Here, we designed a family of stable TIM barrels and explored their stability landscape. The TIM-barrel collection reported in this work exhibits a considerable range in thermostability (more than 60 degrees in \( T_m \)) and conformational stability at 25 °C (more than 25 kcal mol\(^{-1}\) in \( \Delta \Delta G^{25^\circ C} \)). These data can now be used to accelerate the development of future custom design protein stability curves which, in turn, will expand the biomedical and biotechnological applications of \textit{de novo} proteins. For example, by fusion to another \textit{de novo} protein, one of the stabilized scaffolds reported here (NovoTIM13) has been successfully used to create a reaction chamber on the top of the barrel \((56)\), confirming the convenience of working with robust and stable TIM barrels in the path towards functional \textit{de novo} proteins.

In the same way that one explores the sequence space by studying homologous proteins from different organisms, \textit{de novo} design with a fixed backbone follows a similar strategy generating new sequences within the same topology. It is well known that highly stable proteins can be generated by computational design. However, one of the unexpected findings resulting from the thermodynamic characterization of this family of NovoTIMs is that very stable proteins can be obtained in unexplored regions of the stability landscape. The paths followed in the stability landscape of NovoTIMs are severely influenced by epistatic effects that appear to arise from an increase in hydrophobic clusters. The design and characterization of stable \textit{de novo} proteins is an essential step on the route to the next generation of new protein functions charting novel sequence space.

Materials and Methods

**Enzymes and biochemicals**

All reagents were of analytical grade from Merck KGaA®. Water was distilled and deionized.
**Design protocol**

*De novo* TIM barrels were designed using the Rosetta software suite v.3.2 (57, 58; https://www.rosettacommons.org/). All NovoTIMs were designed using NovoTIM0 as template. The script used for the NovoTIM collection follows and executes the steps indicated in supplementary methods and in the algorithm showed in Fig. S1.

**Cloning, overexpression, and protein purification**

The protocol used to clone, overexpress, and purify NovoTIMs is described in detail in supplementary methods.

**Far- and Near-UV circular dichroism**

Circular dichroism (CD) spectra were collected in buffer D: 10 mM NaH$_2$PO$_4$ pH 8.0 in a Chirascan Spectropolarimeter using a Peltier device to control the temperature (Applied Photophysics®). For Far-UV spectra, 0.4 mg mL$^{-1}$ of NovoTIM was used for all measurements (1 nm bandwidth, 185-260 nm wavelength range, 1 mm cuvette). For Near-UV spectra, 1 mg mL$^{-1}$ of NovoTIM was used for all measurements (1 nm bandwidth, 250-350 nm wavelength range, 10 mm cuvette). The spectra for thermally-unfolded states were collected at 90 °C. Spectra for chemically-unfolded states were collected at 9 M urea for all NovoTIMs, except for NovoTIM14, which was collected at 7 M GdnHCl. Raw data were converted to mean residue molar ellipticity ([θ]) using: [θ] = θ/(l C N$_r$), where θ is ellipticity collected in millidegrees, l is the cell path length in mm, C is the NovoTIM molar concentration, and N$_r$ the number of residues per protein. Far-UV spectra were deconvoluted with CDNN (59).

**Intrinsic fluorescence**

Intrinsic fluorescence (IF) spectra were collected on a PC1 ISS Spectrofluorometer (Champaign IL-USA®) equipped with a Peltier device controlling the temperature. In all measurements, protein concentration was 0.4 mg mL$^{-1}$ in buffer D: 10 mM NaH$_2$PO$_4$ pH 8.0 (1 nm bandwidth slits, 295 nm excitation wavelength, 310–450 nm emission wavelength range). Spectra for chemically-unfolded states were collected at 9 M urea for all NovoTIMs, except for NovoTIM14, which was collected at 7 M GdnHCl. Fluorescence spectral center of mass (SCM) was calculated from intensity data ($I_λ$) obtained at different wavelengths ($λ$): SCM = $\sum I_λ/\sum I_λ$.

**Three-dimensional structure determination**

NovoTIM structures were determined by X-ray crystallography as described in supplementary methods. The coordinates and structure factors were deposited in the PDB with accession codes: 6YQY (sTIM11noCys), 6Z2I (NovoTIM6), and 6YQX (NovoTIM13). The figures were created using PyMOL Molecular Graphics System v.2.2.0 (Schrodinger, LLC).

**Analytical size exclusion chromatography**

Hydrodynamic measurements were performed on a Superdex 75 10/300 GL analytical column coupled to an ÄKTA System (GE Healthcare Life Sciences®). All experiments were performed in...
buffer C: 10 mM NaH$_2$PO$_4$ pH 8.0, 150 mM NaCl at 25 °C and a protein concentration range from 0.01 to 2.0 mg mL$^{-1}$. Experimental molecular weight, Stokes-radii, and oligomeric state were calculated from elution volumes and a calibration curve derived from 7 different known proteins.

**Thermal unfolding followed by circular dichroism**

Temperature-induced unfolding was monitored by CD at 222 nm as a function of temperature using 0.4 mg mL$^{-1}$ in buffer D: 10 mM NaH$_2$PO$_4$ pH 8.0, a heating rate of 1.0 and 1.5 K min$^{-1}$, and a 1 mm path-length cell. T-melt experiments were analyzed as indicated in supplementary methods.

**Thermal unfolding followed by differential scanning calorimetry**

Differential scanning calorimetry (DSC) scans were carried out in a VP-Capillary DSC system (MicroCal®, Malvern Panalytical). Samples were prepared by exhaustive dialysis in buffer D: 10 mM NaH$_2$PO$_4$ pH 8.0 and then degassed at room temperature. Corresponding buffer-buffer traces were subtracted from each endotherm. For all proteins a reheating scan was performed to determine the reversibility or irreversibility of the process. To verify that irreversibility was not the result of a too high final scanning temperature, the first scans were also performed heating near the $T_m$. For NovoTIMs with a reversible thermal unfolding, protein concentration varied from 0.2-2.5 mg mL$^{-1}$ and scan rates from 0.5 to 3.0 K min$^{-1}$. For NovoTIMs with an irreversible thermal unfolding, protein concentration was 1 mg mL$^{-1}$ and scan rates from 0.5 to 3.0 K min$^{-1}$. For NovoTIM14 in native conditions, protein concentration was increased to 2.5 and 4.5 mg mL$^{-1}$ to determine accurately the transition. For NovoTIM14 in the presence of urea, all the scans were done at 1 mg mL$^{-1}$ from 2.0 to 6.0 M urea with samples incubated for 6 hours at 10 °C. DSC data were analyzed as indicated in supplementary methods. Origin v.7.0 (OriginLab Corporation, Northampton, MA, USA.) with MicroCal software was used for data analysis.

**Chemical-induced unfolding**

All experiments were carried out at a protein concentration of 0.1 mg mL$^{-1}$ in buffer D: 10 mM NaH$_2$PO$_4$ pH 8.0 at 25 °C. To determine whether urea induced unfolding was reversible, unfolding and refolding experiments were assayed. For unfolding experiments, native NovoTIM was the initial state, whereas for refolding, the starting state was the unfolded NovoTIM incubated overnight in 9.0 M urea. Thereafter samples were incubated at different concentrations of urea (0-9.0 M), either increasing or decreasing the initial concentration (for unfolding and refolding experiments, respectively). Intrinsic fluorescence of both, unfolding and refolding samples, was measured at different times to determine the equilibrium time. Once the equilibrium time was found, unfolding experiments with samples incubated for 12 hours and followed by CD and IF were performed as aforementioned. IF data at fixed emission wavelength and CD data at 222 nm were both collected over 2 minutes at each urea concentration. Chemical unfolding experiments were analyzed as indicated in supplementary methods.

**Stability curve and global thermodynamic stability**

Global stability curves, $\Delta G(T)$, were calculated using the thermodynamic parameters obtained from DSC experiments and the Gibbs-Helmholtz equation (60):
\[ \Delta G(T) = \Delta H \left( 1 - \frac{T}{T_m} \right) - \Delta C_P \left( \frac{T_m - T}{T_m} \right) + T \ln \left( \frac{T}{T_m} \right) \]  

(Eqn. 1)

The area under the stability curve is a measure of the global stability of the protein \((47)\). It was calculated integrating equation 1 from the lowest temperature at which the protein is in the liquid state i.e. 0 °C \((273.15 \text{ K})\) to \(T_m\):

\[ \text{Area} = \left( \Delta H - T_m \Delta C_P \right) \left( T_m - T \right) - \left( \frac{\Delta H}{2T_m} - \frac{\Delta C_P}{2} \right) \left( T_m^2 - T^2 \right) + \frac{\Delta C_P}{4} T_m^2 \left( T^2 \ln \frac{T}{T_m} - \frac{T^2}{2} \right) \]  

(Eqn. 2)

**Stability landscape**

The stability landscape was constructed by plotting \(T_m\) and \(\Delta H\) obtained from thermal unfolding experiments, and \(\Delta G_{25^\circ C}\) obtained from chemical unfolding data. The 3D surface map was calculated using an XYZ gridding approach for randomly spaced data based on the modified Shepard’s method. The expanded matrix was a rectangular array with \(\Delta G_{25^\circ C}\) as Z values whose columns were mapped to \(T_m\) as X values and rows to \(\Delta H\) as Y values. The method constructs a function \(F(x,y)\) go through the experimental data \((T_m, \Delta H,\) and \(\Delta G_{25^\circ C}\)) and interpolates \((F(x,y) = z)\) for all irregular distributed points \((x_i, y_i, z)\). The stability surface was constructed with the software Origin v.7.0 (OriginLab Corporation, Northampton, MA, USA.) and colored according to normalized \(\Delta G_{25^\circ C}\) values in 0.1 bins. It should be noted that although \(T_m\), \(\Delta H\), and \(\Delta G\) are related by equation 1, their surface representation in 3D requires a common \(\Delta C_P\). Therefore, the stability surfaces shown in Fig. 5 and Fig. S14 are not a 3D fitting to the Gibbs-Helmholtz equation.

**Thermodynamic double-mutant cycles**

To calculate non-additive effects between different NovoTIM barrel regions, an approximation based on double mutant cycles was used \((48, 61, 62)\). The thermodynamic cycles were constructed linking single-region/double-region designs and double-region/triple-region designs as indicated in Fig. S12. Each corner of the square represents a different NovoTIM where the mutations are located in a specific region of the barrel or in a combination of them. For double-region cycles (upper panel), from the first to the second design round, \(\Delta G_1\) and \(\Delta G_2\) are the changes in stability produced when a single region of the barrel was mutated. \(\Delta G_1\) and \(\Delta G_2\) are the changes in stability generated when the same mutations are evaluated in the background of another first-round design. In the triple-region cycles (lower panel), from the second to the third design round, \(\Delta G_1\) and \(\Delta G_2\) are the changes in stability produced when the mutations of a single region are introduced in the background of NovoTIM0 or in a double-region design, whereas \(\Delta G_1\) and \(\Delta G_2\) are the changes in stability generated when a double-region design was incorporated in the background of NovoTIM0 or in a single region design, respectively. Considering that \(\Delta G\) is a state property, if two regions of the barrel are energetically-independent, their effects will be additive and not coupled. Therefore, stability changes linked to a particular region will result in the same values on parallel sides of the square, i.e., \(\Delta G_1 = \Delta G_3\) and \(\Delta G_2 = \Delta G_4\). Any difference the values on the parallel sides of the squares indicates a deviation from additivity and measures the coupling energy between different regions of the barrel, given by \(\Delta \Delta G = \Delta G_1 - \Delta G_3 = \Delta G_2 - \Delta G_4\), where \(\Delta \Delta G\) values have been referred as coupling energy, non-additive effects, interaction energies, and more recently epistatic effects \((48)\).
**Sequence and structural analysis**

Sequence alignment was performed with MAFFT v.7.450 (63) using the secondary structure information from the sTIM11 structure (PDB ID: 5BVL). Sequence identity was calculated with the SIAS server (Universidad Complutense de Madrid, 2013). Structural alignments and RMSD calculations were performed using PyMOL Molecular Graphics System v.2.2.0 (Schrodinger, LLC). Cavity volumes were calculated with MOLE v.2.5 (64) using a standard probe radius of 5 Å and an interior threshold of 1.1 Å with a non-directed exploration path. The accessible surface area (ASA) was calculated with VADAR v.1.8 (65). In these analyses, changes in ASA for the unfolded state were calculated with an extended Gly-X-Gly peptide. Hydrogen bonds, as well as salt bridges, were calculated using HBPLUS v.3.06 (66) and ESBRI (67) with default parameters for distances and angles. A salt bridge was assigned when two atoms of opposite charge were observed within 4 Å. Hydrophobic clusters (formed by ILV residues) were calculated following an algorithm previously reported by Sobolev (68) with a script developed by Dr. Noelia Ferruz-Capapey from the Höcker Lab.

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**Competing interests**

Authors declare no competing interests.

**Data and materials availability**

All data to support the conclusions of this manuscript are included in the main manuscript and supplementary information. Coordinates and structure files have been deposited to the Protein Data Bank (PDB) with accession codes: 6YQY (sTIM11noCys), 6Z2I (NovoTIM6), and 6YQX (NovoTIM13).
References


**Figure 1. Modular design approach to obtain the NovoTIM collection.** Cartoon representation of the regions and the corresponding residues modified in each design round. The two cysteine residues present in sTIM11 that were reverted to the corresponding symmetry-related residues in sTIM11nocys are shown in magenta (C8Q and C181V). Mutations W34V and A38G (as well as their 4-fold-symmetry related residues) introduced in NovoTIM0 are shown in black. The inner core, formed by the β-barrel residues A21, R23, I40, I42, A67, R69, I86, and I88 (as well as their 2 fold-symmetry related residues) is shown in orange. The peripheral bottom core, formed by the N-terminal region of even β-strands and the C-region of the flanking α-helices, that is, residues Q11, E15, T18, K31, and V34 (as well as their 4-fold-symmetry related residues) is colored green. Peripheral top core situated at the C-terminal region of the odd β-strands and the N-terminal region of the flanking α-helices formed by residues K2, A5, W6, Y22, S24, and D29 (as well as their 4-fold-symmetry related residues) is shown in purple. All the sequences analyzed in this work are reported in Fig. S2 and tables S1-S2.
Figure 2. Conformational properties and equilibrium unfolding of NovoTIMs. (A) Far-UV CD spectra. (B) Near-UV CD spectra. (C) Intrinsic fluorescence (IF) spectra ($\lambda_{exc} = 295 \text{ nm}$). (D) Thermal unfolding followed by CD$_{222\text{nm}}$ (scan rate: 1.5 K hr$^{-1}$). (E) Differential scanning calorimetry (DSC) endotherms (scan rate: 1.5 K hr$^{-1}$; for easy comparison, the physical and chemical baselines have been subtracted). (F) DSC endotherms of NovoTIM14 in the presence of increasing concentrations of urea (2.0 to 6.0 M) from bottom to top (scan rate: 1.5 K hr$^{-1}$). For clarity, in panels E and F only a small part of the pre- and post-transition baselines are shown. (G) Chemical unfolding using urea and followed by CD (notice that NovoTIM14 does not unfold with urea; error bars indicate the standard deviation). (H) Chemical unfolding induced by guanidinium hydrochloride for NovoTIM14 (squares: CD, circles: IF; error bars indicate the standard deviation).
Figure 3. Stability and energetic coupling in NovoTIMs. (A) Correlation between two parameters which are proportional to the exposed surface area: $m$ value from chemical unfolding and $\Delta C_p$ from temperature-induced unfolding (solid line: linear regression excluding NovoTIM1, NovoTIM11, and NovoTIM12 data; $R^2$: 0.76. Dotted line: correlation reported by 37; error bars indicate the standard error from global fitting). (B) Stability curves calculated from DSC data (lines) using the Gibbs-Helmholtz equation (open symbols show $\Delta G$ values determined by chemical unfolding at 25 °C. Grey dashed line indicates 25 °C). (C) Correlation between the relative global thermodynamic stability (Area/Area$_{NovoTIM0}$) and thermostability ($T_m$) ($R^2$: 0.93). Inset: correlation between $\Delta G$ at 25 °C determined by chemical unfolding and $T_m$ ($R^2$: 0.87). For NovoTIM14, where two transitions were found, it was assumed that the one observed at lower [GdnHCl] corresponds to the lower $T_m$. (D) Thermodynamic cube showing the coupling energy ($\Delta \Delta G_{int}$) between different regions of NovoTIMs. $\Delta \Delta G_{int}$ values were calculated from the double-mutant cycles shown in Fig. S12. $\Delta \Delta G_{int}$ values between single-region mutants are depicted as colored arrows from the top face to the bottom face. $\Delta \Delta G_{int}$ values calculated for the addition of a single-region design to a double-region design are shown as colored arrows in the bottom face.
Figure 4. Three-dimensional structures of NovoTIMs. (A) Structural alignment of X-ray structures of sTIM11 (PDB ID: 5BVL), sTIM11noCys (PDB ID: 6YQY), NovoTIM6 (PDB ID: 6Z2I), and NovoTIM13 (PDB ID: 6YQX). The RMSD $\alpha$ between the structure and the Rosetta model among the quarters in each protein is shown in the lower part of the panel. (B) Comparison of sTIM11noCys and sTIM11 structures (RMSD: 1.07 Å -174 $\alpha$-). The mutated residues 8 and 181 in sTIM11noCys are zoomed in the bottom part. (C) Comparison of the NovoTIM6 structure with the Rosetta model (RMSD: 2.28 Å -168 $\alpha$-). The quarters with the highest and lowest structural similarity are highlighted (bottom left and bottom right, respectively). (D) Comparison of the NovoTIM13 structure with the Rosetta model (RMSD: 1.43 Å -181 $\alpha$-). The quarters with the highest and lowest structural similarity are highlighted (right and left, respectively). Sidechains of the mutated residues are shown in sticks.
Figure 5. Stability landscape of de novo TIM barrels. The stability surface is colored according to normalized ΔG_{25°C} values in 0.1 bins. Colored lines represent a possible pathway from one design to another and were drawn only as a guide to the eye: orange from sTIM11 to sTIM11noCys, cyan from sTIM11noCys to NovoTIM0, yellow from NovoTIM0 to the first-round NovoTIMs, white from single-region designs to the double-region ones, and blue from double-region designs to the triple-region design. T_m and ΔH data were obtained from thermal unfolding, whereas ΔG values derive from chemical unfolding (table 1).
Table 1. Thermodynamic properties of NovoTIMs.

<table>
<thead>
<tr>
<th>de novo TIM barrel</th>
<th>Type of modular design</th>
<th>$I_m$ (ºC)</th>
<th>$\Delta H$ (kcal mol$^{-1}$)</th>
<th>$\Delta C_P$ (kcal mol$^{-1}$ K$^{-1}$)</th>
<th>$\Delta H_{m}/\Delta H$</th>
<th>Global stability (kcal K mol$^{-1}$)$^b$</th>
<th>$\Delta G_{25^\circ C}$ from stability curve (kcal mol$^{-1}$)$^b$</th>
<th>$m$ (kcal mol$^{-1}$ M$^{-1}$)$^c$</th>
<th>$\Delta \Delta G_{int}$ (kcal mol$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>sTIM11</td>
<td>First reported de novo TIM barrel</td>
<td>80.0 ± 0.2</td>
<td>92.9 ± 1.3</td>
<td>2.19 ± 0.19</td>
<td>0.99 ± 0.05</td>
<td>279.6</td>
<td>4.6</td>
<td>1.34 ± 0.06</td>
<td>3.1 N/A</td>
</tr>
<tr>
<td>sTIM11noCys</td>
<td>sTIM11 without cysteines</td>
<td>65.6 ± 0.1</td>
<td>82.0 ± 0.6</td>
<td>2.36 ± 0.08</td>
<td>0.99 ± 0.03</td>
<td>NA</td>
<td>175.8</td>
<td>3.8 ± 0.2</td>
<td>1.9 N/A</td>
</tr>
<tr>
<td>NovoTIM0</td>
<td>Base design</td>
<td>47.0 ± 0.2</td>
<td>24.7 ± 0.6</td>
<td>0.44 ± 0.06</td>
<td>1.09 ± 0.08</td>
<td>NA</td>
<td>60.5</td>
<td>1.5 ± 0.1</td>
<td>2.1 N/A</td>
</tr>
<tr>
<td>NovoTIM1</td>
<td>Single-region design: Internal core</td>
<td>71.0 ± 0.4</td>
<td>48.1 ± 0.8</td>
<td>0.69 ± 0.12</td>
<td>0.98 ± 0.05</td>
<td>NA</td>
<td>225.9</td>
<td>3.8 ± 0.1</td>
<td>1.87 ± 0.10</td>
</tr>
<tr>
<td>NovoTIM6</td>
<td>Single-region design: Bottom cavity</td>
<td>92.3 ± 0.1</td>
<td>124.9 ± 1.5</td>
<td>2.38 ± 0.06</td>
<td>1.03 ± 0.02</td>
<td>NA</td>
<td>542.2</td>
<td>7.9 ± 0.2</td>
<td>1.51 ± 0.08</td>
</tr>
<tr>
<td>NovoTIM8</td>
<td>Single-region design: Top cavity</td>
<td>77.3 ± 0.3</td>
<td>51.9 ± 0.9</td>
<td>1.19 ± 0.10</td>
<td>0.95 ± 0.09</td>
<td>NA</td>
<td>165.2</td>
<td>2.5 ± 0.2</td>
<td>0.85 ± 0.09</td>
</tr>
<tr>
<td>NovoTIM11</td>
<td>Double-region design: Internal core + Bottom cavity (NovoTIM1 + NovoTIM6)</td>
<td>103.5 ± 0.2</td>
<td>61.7 ± 2.4</td>
<td>0.67 ± 0.14</td>
<td>1.02 ± 0.09</td>
<td>NA</td>
<td>523.2</td>
<td>8.4 ± 0.4</td>
<td>1.75 ± 0.08</td>
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<tr>
<td>NovoTIM12</td>
<td>Double-region design: Internal core + Top cavity (NovoTIM1 + NovoTIM8)</td>
<td>108.8 ± 0.3</td>
<td>79.3 ± 1.9</td>
<td>0.72 ± 0.08</td>
<td>1.01 ± 0.08</td>
<td>NA</td>
<td>791.5</td>
<td>10.9 ± 0.2</td>
<td>1.77 ± 0.03</td>
</tr>
<tr>
<td>NovoTIM13</td>
<td>Double-region design: Bottom cavity + Top cavity (NovoTIM6 + NovoTIM8)</td>
<td>92.8 ± 0.4</td>
<td>46.7 ± 4.5</td>
<td>N/D</td>
<td>N/D</td>
<td>120.3</td>
<td>N/D</td>
<td>9.5 ± 0.2</td>
<td>1.54 ± 0.03</td>
</tr>
<tr>
<td>NovoTIM14</td>
<td>Triple-region design: Internal core + Bottom cavity + Top cavity (NovoTIM1 + NovoTIM6 + NovoTIM8)</td>
<td>91.5 ± 0.1</td>
<td>5.4 ± 0.2</td>
<td>N/D</td>
<td>N/D</td>
<td>37.4</td>
<td>N/D</td>
<td>Tot: 26.7 ± 1.7</td>
<td>N to I: 18.0 ± 0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>N to I: 9.3 ± 1.1</td>
<td>I to U: 17.4 ± 0.3</td>
<td>N to I: 1.8 ± 0.2</td>
<td>I to U: 4.4</td>
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<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>N to I: 3.2</td>
<td>I to U: 4.4</td>
</tr>
</tbody>
</table>

$^a$ ± indicate the standard deviation calculated from experiments at different protein concentrations.

$^b$ The global thermodynamic stability was calculated from the area of the stability curve evaluated between 0 ºC and $T_m$.

$^c$ ± indicate the standard error from global fitting.

$^d$ $\Delta \Delta G_{int}$ calculated from NovoTIM11 + NovoTIM8 to NovoTIM14 (see Fig. S12).

$^e$ $\Delta \Delta G_{int}$ calculated from NovoTIM12 + NovoTIM6 to NovoTIM14 (see Fig. S12).

$^f$ $\Delta \Delta G_{int}$ calculated from NovoTIM13 + NovoTIM1 to NovoTIM14 (see Fig. S12).