Folding the proteome

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Protein folding is an essential prerequisite for protein function and hence cell function. Kinetic and thermodynamic studies of small proteins that refold reversibly were essential for developing our current understanding of the fundamentals of protein folding mechanisms. However, we still lack sufficient understanding to accurately predict protein structures from sequences, or the effects of disease-causing mutations. To date, model proteins selected for folding studies represent only a small fraction of the complexity of the proteome and are unlikely to exhibit the breadth of folding mechanisms used in vivo. We are in urgent need of new methods both theoretical and experimental - that can quantify the folding behavior of a truly broad set of proteins under in vivo conditions. Such a shift in focus will provide a more comprehensive framework from which to understand the connections between protein folding, the molecular basis of disease, and cell function and evolution.

Our current understanding of protein folding

Proteins are central to all cellular events: they catalyze chemical reactions, do mechanical work, and perform structural roles in the cell. Although proteins are synthesized as long linear polymers of amino acids, each must fold into a specific 3D structure in order to perform its cellular function. The precise sequence of amino acids has a profound influence on the structure of a protein, and hence its function. Indeed, pioneering experiments by Anfinsen in the 1950s showed that the structure of a protein can be determined exclusively by the sequence of amino acids in the polypeptide chain [1]. Therefore, in theory when provided with the sequence of a new protein we should be able to predict its structure, which can help predict its function. Moreover, a complete understanding of all physical principles that shape protein folding and structure would enable the de novo design of novel protein structures and functions and accurate predictions of the effects of disease-causing mutations on protein structure and folding.

Anfinsen's observations were the wellspring of the protein folding field, and intense effort has been devoted to determining precisely how a full-length polypeptide chain can quickly refold to form its native structure with high efficiency. Similar to Anfinsen, most efforts focus on studying a purified protein that is first chemically denatured to form an enormous ensemble of rapidly interconverting flexible conformations (the unfolded ensemble), then diluted

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away from the denaturant in order to enable the protein to refold and regain its biologically active structure (the native state) (Box 1). These refolding experiments can be used to measure protein folding rates and also protein stability, calculated as the free energy of folding ($\Delta G^{\circ}_{\text{folding}}$) [2]. An important prerequisite for these measurements is the identification of experimental conditions under which measurable concentrations of reactants and products (the unfolded ensemble and native state) exist at equilibrium, and hence is restricted to the subset of proteins that unfold and refold reversibly on an experimentally measurable time scale.

Our focus on proteins that refold reversibly after chemical denaturation means that the subset of proteins whose folding properties are well understood share many structural and folding features. Many of these proteins are small (<100 aa), single-domain, monomeric, marginally stable ($\Delta G^{\circ}_{\text{folding}} \sim -15$ -30 kJ/mol) and fold quickly (ms-s time scale) via concerted mechanisms that lack well-populated intermediate conformations [3–9]. This is not surprising, as it has been shown that proteins that are large, multimeric, and/or fold slowly via long-lived, partially folded intermediate structures are more likely to misfold and aggregate [10–13], and are therefore unsuitable for calculating $\Delta G^{\circ}_{\text{folding}}$ (Box 1).

Detailed studies of proteins that refold reversibly after chemical denaturation have enabled descriptions of many fundamental features of refolding mechanisms, including measurements of typical rates for global hydrophobic collapse and secondary structure formation [14]. Such small proteins are also more amenable to computer simulations than large proteins, and hence continue to provide an invaluable bridge between experimental results and the development of protein folding simulations and theory [15,16]. Our extensive investigations of these model proteins have now revealed common features of their refolding mechanisms [14,17,18], including general correlations between folding rate and the length [5] or structural complexity [3] of the protein, and some common folding intermediates including the 'molten globule', a globally collapsed conformation with significant secondary structure but little or no stable tertiary structure that is often observed shortly after dilution of an unfolded protein from denaturant [19,20]. These and other common mechanistic features identified to date have led to speculation that protein folding in general is now a well-understood phenomenon [21–24]. However, with few notable exceptions [25–28], our current state of knowledge has produced only modest progress towards accurate ab initio predictions of protein structure from sequence, design of novel proteins, or treatments for the molecular basis of protein folding diseases.

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Box 1. The standard set of experiments used to characterize protein folding in vitro

This approach is applicable only to proteins that reversibly unfold and refold in a two-state reaction in vitro. Conditions must be identified that eliminate aggregation (Figure IA), a common side reaction. Protein folding in reversible systems (Figure IB) is often monitored using optical methods including tryptophan fluorescence emission and circular dichroism spectroscopy. Thermodynamic stability $(\Delta G^{\circ}_{folding})$ is measured by equilibrating the protein in various concentrations of a chemical denaturant and measuring the conformation spectroscopically at each denaturant concentration (Figure IB, green). The linear transition region in the sigmoidal unfolding/ refolding titration can be extrapolated to calculate $\Delta G^{\circ}_{\text{folding}}$ in 0 M denaturant. The same spectroscopic tools can be used to monitor refolding and unfolding kinetics (Figure IB, blue). The rate constants for folding $(k_{\rm f})$ and unfolding $(k_{\rm u})$ are dependent on the residual denaturant concentration. By systematically varying the final denaturant concentration, one can construct a V-shaped chevron plot (bottom right) where the folding and unfolding rate constants are plotted versus the final concentration of denaturant [76]. Each 'arm' of the V-shaped plot can be extrapolated to obtain the unfolding and refolding rate constants at 0 M denaturant. For proteins that fold via a simple one-step process, these rate constants can also be used to calculate $\Delta G^{\circ}_{\text{folding}}$

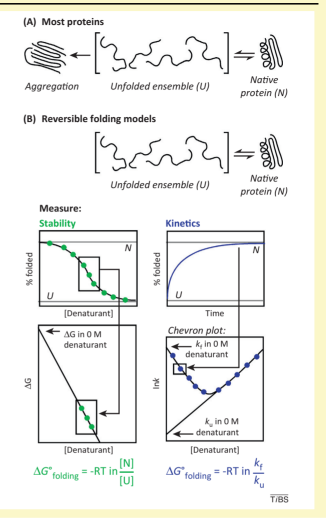
Figure I. (A) For most proteins, dilution of the ensemble of unfolded conformations (U) out of denaturant results in a kinetic competition between correct folding (N) versus aggregation. This results in <100% reversibility of the refolding reaction, meaning that such proteins cannot be used for the thermodynamic and kinetic analyses described below. (B) For proteins that do refold reversibly, there are widespread assays available to study protein stability and folding kinetics. For such assays, conditions must be identified where folding is reversible: upon dilution, the ensemble of unfolded conformations (U) is converted exclusively to the native structure (N) with no detectable aggregation or hysteresis. Under these conditions, the free energy of folding $(\Delta G^{\circ}_{\text{folding}})$ can be determined either from equilibrium denaturation measurements (left; green) or from kinetic measurements of rate constants (k_r and k_{u}) (right; blue). Abbreviations: R, gas constant; T, temperature.

Why is protein folding still a 'problem'?

Why is it that, despite extensive study and the emergence of common features, our ability to predict the folding behavior of a protein from its sequence is still so limited? We wondered whether the intense focus on proteins that unfold and refold reversibly in vitro, which provide valuable tools for building the foundation of our understanding of folding mechanisms, might nevertheless hamper the development of a broader, more comprehensive understanding of protein folding. Unfortunately, the majority of proteins in a proteome cannot refold reversibly after dilution from a chemical denaturant. Instead, once taken out of the cellular environment, chemically denatured and diluted into a buffer designed to mimic physiologically relevant conditions (pH, salt, etc.), most proteins misfold and aggregate (Box 1) [29,30]. Such proteins are rarely used as folding models, despite their widespread appearance in proteomes.

Typical folding models provide incomplete representations of proteomes

Can we extrapolate from our current understanding of protein folding behavior to describe truly general features of protein folding? An accurate extrapolation requires that our model proteins represent the diversity of proteins from



across entire proteomes. To address this question, we compared structural properties of 165 nonredundant proteins commonly used to draw general conclusions regarding protein folding mechanisms [3-9,23] (see Table S1 in the supplementary material online) to properties of the well-characterized Escherichia coli proteome (Figures 1A and 2). This comparison revealed that protein folding models capture only a small subset of protein structural diversity, shared by only 8.4% of the *E. coli* proteome. The diversity of our model proteins therefore does not accurately represent the diversity of the proteome of even a small, simple organism. This discrepancy is most apparent for transmembrane proteins, but proteome diversity is underrepresented even among water-soluble proteins, particularly those that are large and/or multimeric (Figures 1C and 2). Of course, proteins in a proteome can be classified using other properties beyond those shown here, but we consistently observed that current protein folding models fail to adequately sample the diverse properties of the proteome regardless of the metric used. In addition, there is a growing appreciation that the cellular environment introduces additional challenges - including higher temperatures and protein concentrations - not present during in vitro refolding experiments [31,32]. Paradoxically, despite these challenges more proteins fold efficiently in vivo

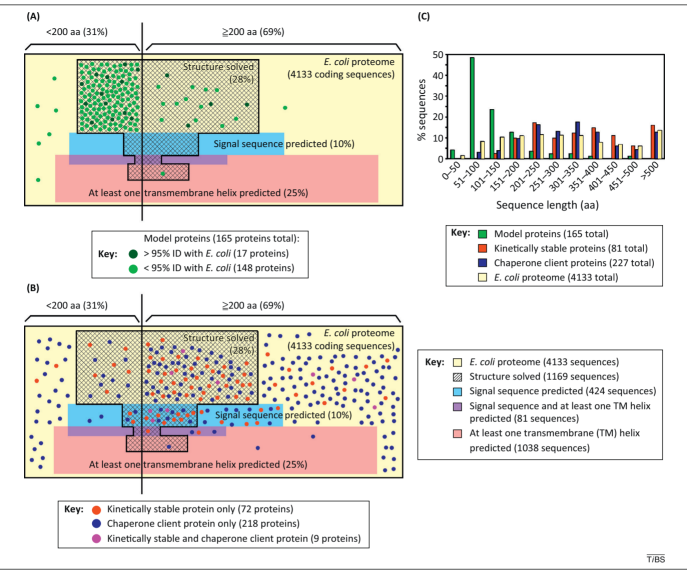


Figure 1. Whereas typical protein folding models exhibit properties not representative of the Escherichia coli proteome, emerging techniques can capture a broader set of proteins. (A) The 4133 proteins from E. coli str. K-12 substr. MG1655 (NC_000913.2) were used to construct a proportional Venn diagram, with each unit area in the yellow rectangle corresponding to one E. coli protein coding sequence. These sequences were divided by length (< or >200 aa) and analyzed for the presence of an N-terminal signal sequence (http://www.cbs.dtu.dk/services/SignalP) (blue shading), one or more transmembrane α-helices (http://www.cbs.dtu.dk/services/TMHMM) (pink shading), both a signal sequence and a transmembrane α-helix (purple shading), and/or a PDB entry with >95% sequence identity to at least some portion of the protein sequence (hatched area). Note that this map underestimates the complexity of the proteome, as each protein coding sequence from the E. coli genome is treated as a separate monomeric protein. A set of 165 nonredundant model proteins used to study protein folding (<95% sequence identity) [3-9] was also analyzed. Each protein is indicated by a green symbol proportional to the size of one E. coli coding sequence. Seventeen of the model proteins have >95% sequence identity to an E. coli protein (dark green symbols); the remaining 148 model proteins are from other organisms (light green symbols). In some cases these models represent individual domains or fragments taken from larger proteins, but as it is known that removal from a larger protein context can change folding behavior [33,44] (see text), the size of the studied domain is used here. (B) Subsets of proteins identified by proteome-wide screens designed to select other, nontraditional folding behavior were categorized as described for the 165 folding models and compared to the properties of the E. coli proteome as in (A). Kinetically stable proteins (red symbols) were identified by protease resistance [43] or resistance to moderate concentrations of sodium dodecyl sulfate (SDS) [42], yielding 81 nonredundant E. coli proteins. E. coli chaperone client proteins (blue symbols) represent both DnaK substrates (category 'enriched' in [61]) and GroEL substrates ('class IV' in [60]), resulting in a set of 227 proteins. Proteins present in both sets (kinetically stable and chaperone client) are indicated as purple symbols. Note that there is only one protein in common between the folding models (A) and kinetically stable and/or chaperone client proteins: maltose binding protein, a kinetically stable protein [43]. (C) Size distribution for each protein group shown in (A) and (B), sorted by sequence length.

than *in vitro* [29,30]. This indicates that the properties that enable folding to proceed efficiently *in vivo* are not necessarily sufficient to enable efficient refolding *in vitro*.

The fundamental physical and chemical principles known to govern protein refolding *in vitro* will of course also apply to larger, more complex proteins in the cellular environment. However, if the protein folding problem was truly solved – if we understood the whole of the physics and chemistry that underlies protein folding – we would now be capable of designing *de novo* a protein structure of any size or complexity, and predicting the effects of disease-causing mutations on protein structure and folding. The fact that we cannot (see e.g., [33]) indicates that our current knowledge is insufficient to solve the problem. In our opinion, closer attention should be paid to proteins currently regarded as unusual outliers as their mechanistic features, which might be considered unusual among our current folding models, could represent footholds for the development of a more comprehensive picture of the diversity of folding mechanisms used across the proteome as a whole.

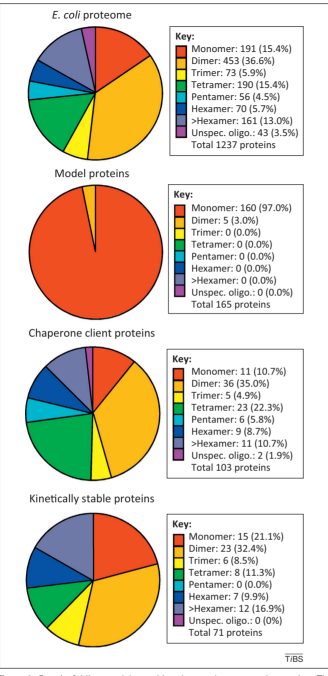


Figure 2. Protein folding models are biased towards monomeric proteins. The multimerization state of each group of proteins shown in Figure 1 (*Escherichia coli* proteome, protein folding models, kinetically stable proteins, chaperone client proteins) was determined. For the *E. coli* proteome, subunit assignments in the Uniprot database were used (30% of proteins in the *E. coli* proteome have assignments; 1237 proteins). Multimerization state for the 165 nonredundant protein folding models was assigned based on reported multimerization state in the protein folding literature. The multimerization state is indicated for 71 of the 81 kinetically stable proteins was assigned using the Uniprot database. Out of the 227 nonredundant chaperone client proteins, 103 have a subunit assignment in the Uniprot database.

The purpose of this article is to highlight five common perceptions regarding protein folding mechanisms developed from studies of small model proteins that in our opinion are unlikely to scale to the folding behavior of all proteins, with the intent to inspire new technology development and research approaches – both *in vitro* and *in vivo* – to close these gaps and increase the predictive power of protein folding research.

Do most proteins globally unfold/refold during their lifetime?

For reasons described above, there has been a strong emphasis on the study of model proteins that unfold and refold reversibly. These proteins tend to be only marginally stable and have fast unfolding and refolding kinetics. These features could mean that these proteins will populate the unfolded state several times over their lifetime in the cell (Figure 3A). The marginal stability and fast kinetics of most model proteins have focused considerable attention on the structural properties of proteins in their chemically denatured ensemble as a model for their unfolded state [34] and the transition between unfolded and native states. However, most proteins do not refold reversibly. Instead, global (or even partial) unfolding can lead to misfolding and aggregation (and/or degradation in vivo) [35]. This suggests that proteins in general will employ strategies to avoid unfolding over their lifetime in the cell. Protein unfolding can be avoided by increasing protein stability, but this tends to rigidify the native structure, which can reduce flexibility needed for binding and catalysis. An alternative option is to increase the energy barrier between the native and denatured states, creating proteins

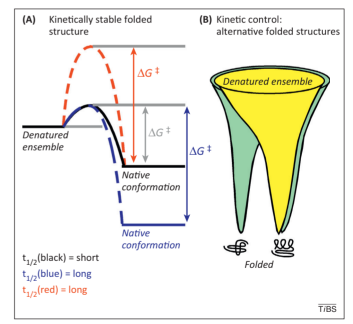


Figure 3. Examples of diversity among protein folding mechanisms. (A) Most proteins currently used as folding models are marginally stable (black), meaning that their folded lifetime $(t_{1/2})$ is short. Lifetime can be increased in two ways. The native structure can be stabilized thermodynamically, increasing the energetic difference between the denatured ensemble and the native structure (increasing $\Delta G^{\circ}_{\text{folding}}$, blue). Alternatively, the energetic barrier separating the denatured ensemble and the native conformation (ΔG^{\ddagger}) can be increased (red); this will preserve the (low) thermodynamic stability but increase the folded state lifetime. Increasing the energy barrier yields kinetically stable proteins, which can be identified by proteome-wide folding screens [42,43] (Figure 1B). (B) Proteins fold from an ensemble of unfolded states, represented by the wide top of a protein folding funnel. In simple model systems (yellow), the funnel has one energy minimum, the native conformation. However, some proteins have a more complex energy landscape and can adopt alternative folded structures (green). These two folded structures may interconvert, or features of the cellular environment may stabilize a subset of early folding intermediates, resulting in a biased accumulation of one structure versus the other(s).

that are kinetically stable but thermodynamically marginally stable, or even less stable than the unfolded state (Figure 3A). Transthyretin [36], P22 tailspike [37], and GFP [38] are three well-studied examples of proteins with high kinetic stability. In vitro, each of these proteins exhibits pronounced hysteresis and hence does not refold reversibly from a chemically denatured state. However, all three fold efficiently in vivo, indicating the cellular environment can alter the energy landscape for folding to arrive at a kinetically trapped native structure. Detailed investigations into the folding mechanisms of these proteins have provided valuable insights into the kinetic competition between folding versus aggregation, including uncovering the effects of conformational changes within a single polypeptide chain on aggregation propensity [39], introducing the concept of folding mutations that can alter partitioning between folding and aggregation without altering the stability of the native state [40] and highlighting the role of folding during translation as a mechanism to increase folding yield in vivo [41].

Many kinetically stable proteins are resistant to unfolding in the presence of moderate concentrations of sodium dodecyl sulfate (SDS) [42] and/or to protease digestion [43]. Proteome-wide screens developed to identify proteins with these characteristics have revealed that kinetically stable proteins sample some properties of the proteome more effectively than typical folding models (Figures 1B, 2, and Table S2 in the supplementary material online) [42,43]. Indeed, these screens underrepresent small (<200 aa) proteins, which might reflect a lower propensity for kinetic stability among small proteins or a bias within the screen against smaller proteins, similar to technical limitations that impose a bias against transmembrane proteins. Traditionally, kinetically stable proteins are rarely selected as folding models because refolding from a chemically denatured state in vitro can require these proteins to populate long-lived, partially folded conformations that may be prone to aggregation. We suggest that focusing our efforts to understanding the structural features that confer kinetic stability, and the mechanisms used to populate such native structures *in vivo*, provides an opportunity to explore a folding strategy used by a diverse set of proteins that can reduce aggregation *in vivo*.

Do larger, multidomain proteins fold like their component domains?

The vast majority of proteins used for folding studies are both monomeric and consist of a single structural domain (Figures 2 and 4). By contrast, larger proteins often consist of multiple domains. This modular architecture makes it tempting to hypothesize that larger proteins will fold via a hierarchical mechanism whereby local contacts first determine the folding properties of individual domains, followed by the formation of interdomain interactions. This model assumes that the complexities of multidomain protein folding can be reduced to the independent folding of the constituent domains. Whereas such behavior has been observed for some proteins, for others the placement of a domain within the context of a larger multidomain protein can significantly alter the energy landscape for folding [44], making it impossible to predict the effects of disease-causing mutations or other sequence alterations from studies of isolated domains alone [33]. For example, whereas yeast phosphoglycerate kinase (PGK) can be studied as two separate, independently folding domains – both of which appear in the list of 165 model proteins (see Table S1 in the supplementary material online) - the E. coli PGK homolog has a nearly identical structure but unfolds five orders of magnitude more slowly, and its C-terminal domain cannot adopt its native structure in isolation [45]. Even for proteins composed of seemingly 'independent' domains connected by flexible linkers, contextual effects can alter folding efficiency. For example, Ig domains within titin form a beads-on-a-string architecture that was shaped by gene duplication events. Concomitant folding of two adjacent titin domains increases the likelihood of non-native interdomain interactions akin to domain swapping, which occur faster than the folding rate and therefore retard accumulation of the native structure [46]. Similar non-native interdomain interactions reduce the folding rate for calmodulin [47]. Interestingly, evolution has reduced the sequence identity between adjacent titin domains, which reduces the likelihood of misfolding [44].

A related question is the extent to which protein folding thermodynamics and kinetics are affected by the assembly of subunits into a multimeric protein. Studies of intrinsically disordered proteins have shown that folding and binding can be interdependent processes [48,49]. It will be valuable to adapt these approaches to study more subtle effects of subunit interactions on the thermodynamics that govern the folding of multimeric proteins with ordered subunits. More broadly, the complex and crowded environment of the cell provides numerous opportunities for nonspecific binding events not captured during refolding of a purified protein in vitro [50], but our understanding of how protein sequences might have evolved to minimize or modulate these non-native interactions is still in its infancy. We predict that negative design strategies such as these - strategies that suppress aggregation and unproductive nonspecific interactions, rather than stabilize the native structure – will play an increasingly important role as we move towards the *de novo* design of larger, multidomain and multimeric proteins.

Do most proteins populate only a single native structure?

Anfinsen showed that RNase can refold to its native, active protein structure after chemical denaturation, presumably because this structure represents the global energy minimum on a funnel-like energy landscape (Figure 3B, yellow). Most folding models share this feature of a thermodynamically controlled folding pathway. But there are now many examples of proteins with multiple, distinct 'native' conformations separated by large energy barriers [51,52] (Figure 3B, green), including α -lytic protease, whose folding properties are well characterized [53] but which is rarely included in general analyses of protein folding behavior. For proteins such as α -lytic protease that fold under kinetic control, the conversion between these alternative folded structures is often an essential, regulated component of the functional cycle of these proteins [54-56]. Although it is not yet known what fraction of the proteome can populate two or more alternative folded

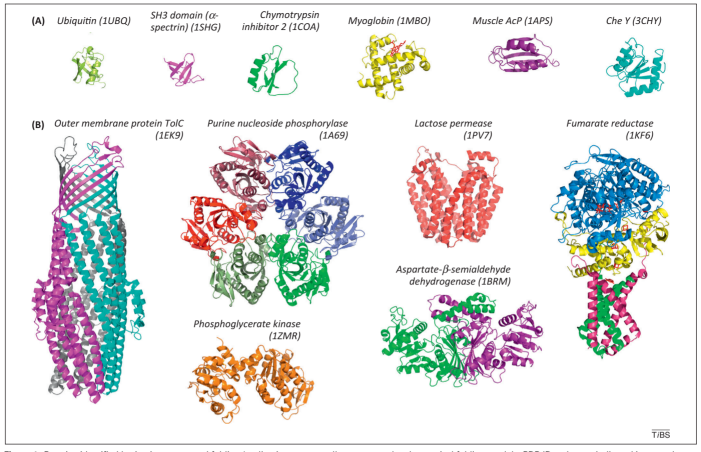


Figure 4. Proteins identified by *in vivo* assays and folding 'outliers' are structurally more complex than typical folding models. PDB ID codes are indicated in parentheses. Subunits of multimeric proteins are shown in different colors, and cofactors (in myoglobin and fumarate reductase) are shown in red. Most models used to study protein folding (**A**) are smaller and less complex than proteins representing diverse properties from the *Escherichia coli* proteome (**B**). Of the *E. coli* proteins shown here, purine nucleoside phosphorylase (a hexamer) and phosphoglycerate kinase (a monomer) were identified in the screen for kinetically stable proteins [43], aspartate- β -semialdehyde dehydrogenase (a dimer) was identified in the screen for chaperone client proteins [60], and the outer membrane protein ToIC (a trimer) was identified in screens for both kinetic stability and chaperone clients [42,61]. Lactose permease (a monomer) is an α -helical transmembrane protein. Two subunits of the tetrameric fumarate reductase (shown in green and pink). The soluble subunits of fumarate reductase (shown in blue and yellow) were identified in the screens for chaperone clients [60,61].

states, our tendency to avoid selecting such proteins as folding models hampers their discovery and hence our understanding of the mechanisms that underlie these folding pathways.

To what extent do molecular chaperones affect protein folding in vivo?

In the cell, proteins fold in an environment that includes molecular chaperones, a class of proteins that has evolved to facilitate the folding and/or suppress the aggregation of other proteins [57-59]. When these proteins were first identified, it was initially thought that chaperones - some of which are essential for cell viability – would explain the higher efficiency of protein folding in vivo versus in vitro. More recently however, quantitative proteomics have revealed that the substrate repertoire for molecular chaperones is fairly narrow under normal growth conditions [60-62], and hence cannot fully explain the higher efficiency of folding in vivo. Yet, although chaperones contribute to the productive folding of only 10-20% of the proteome, their substrates represent a much more diverse subset of proteins than current folding models (Figure 1B and Table S3 in the supplementary material online). These obligate chaperone substrates therefore provide an excellent opportunity to explore the features of folding mechanisms and native structures that lead to chaperone recruitment *in vivo*, versus those that do not. However, because these proteins are unlikely to refold reversibly we will need alternative approaches to study such mechanisms, for example, by exploiting experiments that can quantify the kinetic partitioning between folding and aggregation, or by developing single-molecule experiments that enable a protein to fold at or near infinite dilution, thereby reducing intermolecular interactions between substrate proteins that can lead to aggregation [63,64].

Do other aspects of the cellular environment affect protein folding in vivo?

Our growing appreciation of the contributions of the cellular environment to protein folding has inspired the development of exciting new experimental approaches, including specific fluorescence and isotope labeling strategies that can explore the effects of this complex environment on protein folding thermodynamics and kinetics [65–67]. To date, many of these studies have relied on the same model proteins favored for *in vitro* refolding experiments. Perhaps not surprisingly, in some cases the cellular environment has minimal effects on the folding of these proteins, versus

refolding in vitro; these proteins were selected as folding models in part due to their ability to fold robustly under a wide range of conditions. What is less clear is how other, more diverse types of proteins might exploit properties of the cellular environment to modulate their energy landscape for folding and increase folding efficiency. For example, the folding mechanism for a very small protein might not be altered by folding vectorially (from N to C terminus) during translation or secretion. By contrast, longer proteins take longer to synthesize – and secrete – and therefore spend more time with the N-terminal portion of the protein available for folding prior to the appearance of the C terminus. Indeed, several larger proteins are now known to have cotranslational folding mechanisms in which the energy landscape for folding is significantly altered versus refolding in vitro, leading to the population of folding intermediates not detected during refolding in vitro. These differences often include significant amounts of native-like structure formation for the most N-terminal portions of the nascent protein chain even after tens or hundreds of C-terminal residues have been added to the growing polypeptide chain [41,68,69]. Ideally, computer simulations would enable modeling of co-translational folding processes and the formulation of experimentally testable hypotheses regarding these altered energy landscapes. Unfortunately, the complexities of both longer proteins and the cellular environment greatly increase the complexity of computer simulations, although progress is being made in these directions [70,71].

Concluding remarks and future perspectives

Similar to many biological mechanisms, protein folding is a complex, multifaceted process. *In vitro* refolding studies of small proteins have played an invaluable role in developing our understanding of many fundamental aspects of this process. However, the folding problem is not yet solved, and the technical constraints of any one approach will render it insufficient to fully understand such a complex process. Hence, in our opinion the field of protein folding is currently 'mature' only in the sense that we have extracted much of the information available from our established methods and model systems. A comprehensive understanding of protein folding, particularly in the cell, will require addressing additional phenomena that are not amenable to detection using our current models and approaches.

For these reasons, we advise extreme caution before concluding that the folding behavior of a protein appears to be an unusual outlier. Proteomes contain thousands of proteins, and to date only a small fraction of their diversity has been explored through folding studies. Given that our selection of model proteins to date has been far from general, it remains an open question to what extent unusual 'outlier' folding mechanisms contribute to the folding properties of the proteome as a whole. Studying more diverse proteins, such as those identified from proteomewide screens that do not require reversible refolding (Figures 1B and 4), would enable us to build upon the fundamental laws of physics and chemistry that form the foundation of our current understanding of protein folding in vitro, and learn how the cellular environment exploits those laws in order to build functional cells.

We predict an exciting revolution on the horizon, as the field of protein folding develops the new experimental approaches needed to describe the folding properties of an increasingly diverse set of model proteins and explore the influence of the cellular environment on these proteins. In addition to the examples described above, we are heartened by the recent development of new techniques to study the folding features of transmembrane proteins [72–75], which represent a significant fraction of all proteomes (Figure 1) but by and large have been refractory to traditional approaches to study protein refolding *in vitro*. Once new methods and model systems are in place, we will be able to explore the folding of all corners of the proteome and develop a truly general understanding of how proteins fold.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tibs.2013. 05.001.

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