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Structure of the transition state for the binding of c-Myb and KIX highlights an unexpected order for a disordered system

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A classical dogma of molecular biology dictates that the 3D structure of a protein is necessary for its function. However, a considerable fraction of the human proteome, although functional, does not adopt a defined folded state under physiological conditions. These intrinsically disordered proteins tend to fold upon binding to their partners with a molecular mechanism that is elusive to experimental characterization. Indeed, although many hypotheses have been put forward, the functional role (if any) of disorder in these intrinsically denatured systems is still shrouded in mystery. Here, we characterize the structure of the transition state of the binding-induced folding in the reaction between the KIX domain of the CREB-binding protein and the transactivation domain of c-Myb. The analysis, based on the characterization of a series of conservative site-directed mutants, reveals a very high content of native-like structure in the transition state and indicates that the recognition between KIX and c-Myb is geometrically precise. The implications of our results in the light of previous work on intrinsically unstructured systems are discussed.

kinetics | mutagenesis | protein folding

In the fourth century B.C., Aristotle claimed: “A thing is more properly said to be when it exists in actuality than when it exists potentially.” For example, a log of wood is potentially many objects but becomes a table only once given its proper shape by an experienced carpenter. Because the form is what sculpts a thing to be what it is, the nature of things lies in their form. Consequently, one of the most intuitive concepts that we develop from childhood is that shape determines function. In the context of the intricate environment of the cell, the specificity that characterizes the interactions between different proteins was generally explained by invoking a precise geometrical complementarity between the interacting partners. Thus, the recent proposal that up to 40% of the human proteome appears to be essentially disordered highlights a conundrum (1). Is there a potential value for a protein to be disordered? Given that a disordered protein is more vulnerable, is there a functional advantage in being disordered?

A problem of interest is how to reconcile the accepted structure–function dogma with the existence of disorder (2, 3). Because intrinsically disordered proteins (IDPs) exist but are very elusive to characterization, relevant experimental work is still relatively scarce and such hypotheses remain mostly untested. From a thermodynamic perspective, it has been suggested that destabilization of the native protein structure, leading to its unfolding, lowers the affinity of a protein for its ligand, without necessarily abolishing specificity (4–6). Furthermore, the highly dynamic nature of IDPs was suggested to promote alternative binding of the same disordered segments to different partners, the so-called “moonlighting effect” (7, 8), increasing the repertoire of activities. An interesting mechanistic model, proposed by Wolynes and coworkers (9), suggests that the IDPs display an increased capture radius to recruit and bind partners. According to this hypothesis, commonly referred to as the fly-casting mechanism,

a disordered protein should form with its physiological partner a high-energy complex that would be locked in place by the coupled folding reaction. An important corollary of the fly-casting mechanism is that a potential advantage for a protein to be disordered is to increase the probability (and speed) to bind the interacting partner, due to its extended conformation. Importantly, the fly-casting was shown to become more effective when the unfolding free-energy barriers of proteins are relatively small (10), which implies an anticorrelation between the binding rate constant and protein stability.

The CREB-binding protein (CBP) is a coactivator that modulates the interaction between DNA-bound activator proteins and the components of the basal transcription complex. A globular domain of CBP, namely the KIX domain, is the principal mediator of such interactions (11). Despite its small size, 87 amino acids, and a relatively simple fold, the KIX domain binds different IDP systems via two distinct, but energetically connected, binding sites, called “c-Myb” and “MLL” sites (named after two characteristic ligands of each site, i.e., the transactivation domain of the protein c-Myb and the mixed lineage leukemia, MLL, protein) (12). We have recently characterized the recognition mechanism by which the KIX domain binds to the transactivation domain of c-Myb (13), an IDP that acquires a helical structure when bound to its partner (14). A complete analysis of temperature jump and stopped-flow kinetics in the presence and in the absence of 1,1,1-trifluoro-ethanol, which stabilizes the folded state of c-Myb, revealed that this IDP system recognizes KIX by

Significance

A considerable fraction of proteins, although functional, does not display a well-ordered native state, posing the structure–function dogma into question. Although different models have been described, the role of protein disorder is still shrouded in mystery. Here, we investigate the mechanism of recognition between an intrinsically unstructured protein and its partner. Unexpectedly, we found the binding to occur with a very high degree of geometrical precision, suggesting that a potential value of disorder is not, in this case, to be searched in the speeding up of the reaction thanks to an increased capture radius. Given that in the cell disordered polypeptides may be more vulnerable than folded domains, the advantage in being natively unfolded remains a conundrum.

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following a folding-after-binding scenario, whereby it binds in a relatively unstructured conformation, with the locking of the hydrogen bonds of the helical structure occurring only downhill the primary rate-limiting step.

In this work, we present a detailed description of the structural features of the transition state for the binding-induced folding reaction of c-Myb to KIX. The transition state structure was unveiled using the Φ -value analysis (15, 16). By this technique, residue-specific structural information is inferred by comparing the kinetics of the reaction (folding and/or binding) of the wild-type protein with a series of conservative single mutants, yielding the so-called Φ -value that represents an index of native-like structure of the mutated residue in the transition state. Our results reveal that the transition state of c-Myb is well ordered, with an average Φ -value of 0.89. This is consistently higher than what is generally observed for the folding of single-domain proteins (typically displaying an average Φ -value of 0.36), and indicates that the interaction between KIX and c-Myb is characterized by a considerable geometrical fit.

Results and Discussion

Experimental Design and Strategy of Mutagenesis. Because of their intrinsic nature, transition states never accumulate; thus, their structure must be inferred indirectly. By systematically mutating side chains while probing the effect of the perturbation on the activation- and ground states' free energies, it is possible to map interaction pattern(s) in metastable states and to unveil their structure (15, 16). In practice, the analysis is performed by measuring the so-called Φ -value, which is formalized as

$$\Phi = \frac{\Delta\Delta G_{\#}^{wt-mut}}{\Delta\Delta G_{Eq}^{wt-mut}} \quad [1]$$

Thus, the Φ -value is an index that directly reports on the extent of native-like structure in the transition state, approaching 1 and 0 for native- and denatured-like structures, respectively. As detailed below, we applied this methodology to the IDP system c-Myb in its interaction with KIX.

The aim of the mutational analysis is therefore to obtain structural information on c-Myb in the transition state. With the exception of position 21, which is a glycine residue, all positions of c-Myb were mutated. The substitution of an alanine for a glycine typically stabilizes a helix by 0.4–2 kcal mol⁻¹ (17) by means of a complex mechanism, possibly implying multiple factors including differences in backbone conformational entropy in the denatured state, burial of hydrophobic surfaces on folding, and disruption of hydrogen bonding between the protein and the solvent (18). Thus, Ala–Gly scanning at the solvent exposed sites within the helices is an accepted method to monitor the formation of secondary structure in protein folding as exemplified in ref. 19. Therefore, positions 3, 7, 10, 14, 17, 18, and 22 were mutated to both Ala and Gly. In total, 31 mutants were produced, expressed, and characterized (listed in Table 1). Two of the mutants (L8A and L12A) abolished the binding to KIX and were thus excluded from the kinetic analysis. The remaining mutants were subjected to binding-induced folding experiments and, as described below, 20 reliable Φ -values could be calculated.

Binding Kinetics of Wild-Type c-Myb and Its Site-Directed Mutants.

The binding of c-Myb to KIX is, in theory, a complex reaction involving at least two steps, the folding of c-Myb and a recognition event (14). In a previous study we showed that, despite such a complexity, the observed kinetics appears to follow a two-state mechanism without the accumulation of intermediates (13). An analysis of the experiments obtained in the presence and in the absence of 1,1,1-trifluoro-ethanol, which stabilizes the folded state of c-Myb, revealed complex formation to take place via

a folding-after-binding scenario. Accordingly, c-Myb recognizes KIX in a relatively unstructured conformation and the locking of the hydrogen bonds of the main chain occurs only downhill the rate-limiting step. In an effort to provide a structural interpretation of the reaction, we resorted to characterizing the transition state by Φ -value analysis. In particular, wild-type c-Myb and its site directed mutants were analyzed by independent experiments aimed at the measurement of the association and dissociation rate constants.

We have previously shown that an improved signal-to-noise ratio of the fluorescence change associated with binding can be achieved by using an engineered construct containing a Trp in position 72 of the KIX domain, leading to the variant pwtKIX, and a fusion tag protein at the N-terminal region of c-Myb, namely the construct c-Myb*. Independent control experiments indicated that both the presence of Trp in KIX and of the tag in c-Myb had little effect on the observed binding mechanism and dissociation constant; thus these constructs have been used for the Φ -value analysis. We performed pseudo-first-order binding experiments using c-Myb* and the different site-directed mutants by mixing a constant concentration of pwtKIX (5–10 μ M) with increasing concentrations of c-Myb*, typically ranging from 20 to 100 μ M. Under all conditions, observed kinetics was consistent with a single-exponential behavior, suggesting the reaction follows a two-state mechanism, without the accumulation of intermediates. The observed rate constant for complex formation between KIX and c-Myb* and its mutants as a function of concentrations (Fig. 1) follows bimolecular kinetics. Given the pseudo-first-order approximation, the association rate constants were obtained from the slope of the dependence of the observed rate constant on [c-Myb]. The calculated values are reported in Table 1.

Analysis of the data in Fig. 1 allows one, in theory, to calculate the dissociation rate constant by extrapolating the observed rate constants to zero concentration. Very often, however, the experimental error arising from this extrapolation is too high; therefore, accurate determination of dissociation rate constants may demand a different approach. In analogy to classical experiments on myoglobin (20), a powerful method to measure the dissociation rate is to carry out displacement kinetic experiments where a preincubated complex between two partners is challenged with an excess of a competing reactant. In this work, we performed displacement kinetic experiments where a complex between pwtKIX and c-Myb* was rapidly mixed with an excess of wild-type KIX (i.e., the wild-type domain with a Tyr at position 72). The dissociation time course was then measured at different relative concentrations of wild-type KIX, ranging from 2- to 10-fold, and the observed rate constants were found to be insensitive to the concentration for all mutants. The measured dissociation rate constants for wild-type c-Myb and its site-directed mutants are also listed in Table 1.

Φ -Value Analysis and the Structure of the Folding Transition State.

Following Fersht et al. (15), Φ -values were calculated by dividing the effect of mutation on the activation free energy by the change in free energy of the ground state (as formalized in Eq. 1). For each mutant, the changes in free energies were calculated by applying the following equations:

$$\Delta\Delta G_{\#}^{wt-mut} = RT \ln \left(\frac{k_{on}^{mut}}{k_{on}^{wt}} \right) \quad [2]$$

and

$$\Delta\Delta G_{Eq}^{wt-mut} = RT \ln \left(\frac{K_D^{wt}}{K_D^{mut}} \right). \quad [3]$$

Because of experimental uncertainties (21), Φ -values arising from mutations whose $\Delta\Delta G_{Eq}^{wt-mut}$ was lower than 0.4 kcal mol⁻¹

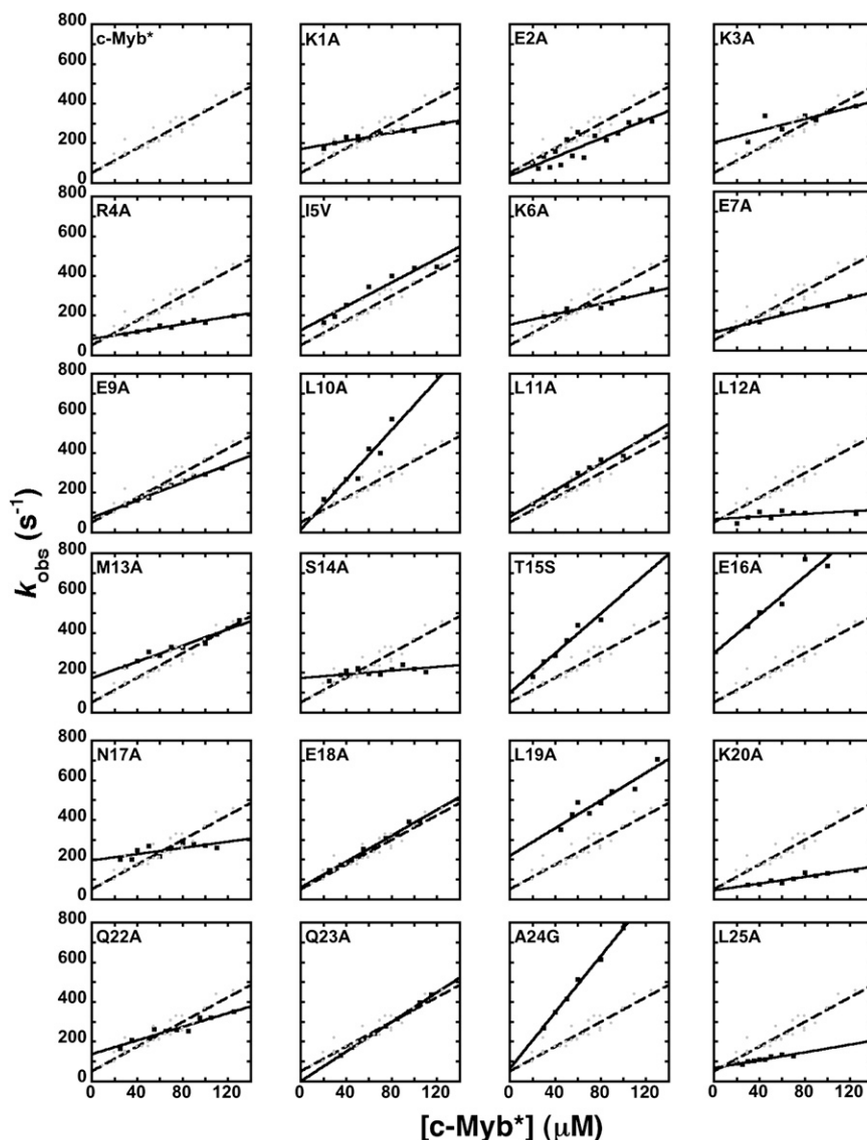


Fig. 1. Pseudo-first-order kinetics of the binding between pwtKIX and c-Myb* at pH 7.2 and 10 °C. Data were recorded at a constant concentration of pwtKIX, typically between 5 and 10 μM , mixed with variable concentrations of c-Myb* and its site-directed mutants. The linear concentration dependence for wild-type c-Myb* is shown for comparison throughout, depicted in gray dots and broken lines. Observed time courses were consistent with single-exponential behavior in all cases.

LFERs were originally introduced to assess the position of the transition state along the reaction coordinate during the formation of a covalent bond (24); in particular, by altering the structure and thus the reactivity of a substrate, the dependence of activation free energy on ground-state free energy generally yields a linear profile. The slope of the observed correlation, classically denoted as α , reflects the position of the transition state along the reaction coordinate. Surprisingly, LFER plots not only apply to simple organic reactions but also to complex reactions stabilized by many noncovalent interactions, and have been then used in enzymology (25), binding reactions involving allosteric control (26, 27), and protein folding (28).

In the case of protein folding, it has been suggested that the linearity of the LFER plot is distinctive of the so-called nucleation-condensation mechanism, whereby the transition state resembles a distorted version of the native state and the whole protein collapses around a weakly formed nucleus (29). Comparative analysis of nearly all Φ -values reported to date revealed that the α -value in protein folding is robust and, for nearly all

proteins investigated, α is about 0.36 (30). The mutational work reported in Table 1 is an opportunity to apply the LFER to the recognition reaction between an IDP and its physiological partner (Fig. 3). In agreement with the behavior recalled above for protein folding, also the folding-after-binding reaction of c-Myb* to KIX displays a linear LFER. Unexpectedly, however, the calculated α -value is 0.89, which suggests the transition state to display a very high degree of native-like structure. Therefore, the rate-limiting transition state does not display an increased capture radius of c-Myb*, and it appears to be more ordered than what is typically observed for the folding reaction of globular proteins. These observations allow us to conclude that, despite being an IDP, c-Myb is characterized in its interaction with pwtKIX by a high degree of geometrical precision, and the protein appears by-and-large folded in the transition state. This is in stark contrast with what was recently observed on another IDP system by Jemth and coworkers, who observed an α of about 0.24 for the binding-induced folding of the two proteins NCBD and ACTR (31).

