Relative Solvent Accessible Surface Area Predicts Protein Conformational Changes upon Binding

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DOI 10.1016/j.str.2011.03.010

INTRODUCTION

Interactions between polypeptide chains are integral to most biological processes. Our understanding of the molecular mechanisms underlying these interactions has been greatly enhanced by the determination of a large number of three-dimensional structures of proteins in both monomeric and complexed states. This has revealed that protein interactions can be associated with varying amounts of conformational change, from slight shifts in the conformations of side chains to large changes in the relative orientations of domains (Gerstein et al., 1994; Betts and Sternberg, 1999; Janin et al., 2007). Protein interactions can also be accompanied by substantial disorder-to-order transitions in the case of intrinsically disordered proteins (IDPs), which are disordered in isolation but which can often be induced to fold in the presence of binding partners (Wright and Dyson, 2009).

Previously, others have noted a relationship between the conformational changes a protein undergoes upon binding and the interface size of the resulting complex. Janin et al. predicted that complexes with large interfaces would require major structural changes upon binding due to the excessive accessible surface of their isolated subunits (Janin et al., 1988). Later, it was observed that the subunits of protein complexes with large interfaces (>2000 Å²) tend to undergo greater conformational changes upon binding than those with smaller interfaces (Lo Conte et al., 1999). Large interfaces relative to protein size have also been noted as a feature of IDPs that fold only upon binding (Gunasekaran et al., 2003, 2004; Levy et al., 2004; Mézáros et al., 2007). However, despite the recognition of this relationship, there have been few analyses of how conformational changes correlate with interface size or other structural features.

The connection between the intrinsic flexibility of unbound proteins and the conformations they adopt upon binding has been the subject of much speculation. In particular, the conformational selection model has gained prominence. This postulates that proteins exist as ensembles of thermally accessible conformations while free in solution, and that binding occurs via transiently formed conformations that resemble the bound state (Tsai et al., 1999). Computational and experimental studies have provided strong evidence that bound-state-like conformations often exist within the free-state ensemble (Boehr et al., 2009; Gsponer et al., 2008; Lange et al., 2008; Marsh et al., 2010; Tobi and Bahar, 2005). While there is some evidence that more flexible proteins tend to undergo larger conformational changes upon binding (Dobbins et al., 2008), little difference was seen between the predicted flexibility of protein binding sites that undergo large conformational changes upon ligand binding and those that do not (Gunasekaran and Nussinov, 2007).

Much work has focused upon trying to predict the structures of protein complexes given the known structures of free subunits, i.e., docking. While current methods have demonstrated a high degree of accuracy for predicting complexes when the subunits maintain conformations close to their free states, docking is much more difficult for proteins that undergo significant conformational changes. Nevertheless, recent approaches incorporating protein flexibility have considerably improved the accuracy of docking (Andrusier et al., 2008; Bonvin, 2006; Wang et al., 2007).

In this study, we start with an approach that may be considered the opposite of docking: given the known structure of a protein complex, we seek to predict the extent of conformational change the constituent subunits undergo upon binding. We show that, while interface size shows some correlation with conformational change, the solvent accessible surface area of a bound subunit relative to the value expected for a monomeric protein of its size is a much better predictor. By applying this method to a
large number of protein complexes, we observe that proteins tend to undergo larger conformational changes than would be expected from only considering cases where both monomeric and complex structures are available. In addition, we observe a significant increase in intrinsic disorder in regions predicted to undergo large conformational changes upon binding. Finally, we show that the relative solvent accessible surface area of a bound subunit can be used to predict the likelihood that it is highly flexible or disordered in its monomeric state. Even more interestingly, we find that the relative solvent accessible surface area of monomeric proteins is also useful for predicting their intrinsic flexibility and their expected conformational changes upon binding. This demonstrates a strong relationship between protein flexibility and binding and has immediate practical implications for the modeling of protein complexes.

RESULTS AND DISCUSSION

Interface Size Shows Limited Correlation with Conformational Changes upon Binding

We searched the Protein Data Bank (PDB) (Berman et al., 2000) for nonredundant pairs of structures where the same or nearly identical polypeptide chains are observed in monomeric and complexed states (see Experimental Procedures). In total, we collected 68 pairs of matching monomer/homomers and 117 pairs of matching monomer/heteromers. Note that since monomeric structures exist, these are transient (as opposed to obligate/permanent complexes, essentially by definition) (Nooren and Thornton, 2003).

In order to assess the relationship between interface size and binding-induced conformational changes, we plotted interface size versus the all-atom root mean square deviation (rmsd) for the full polypeptide chains between bound and unbound conformations for all pairs in the monomer/complex set (Figure 1A). There are only moderate correlations between interface size and log(rmsd) for homomers \((r = 0.59)\) and heteromers \((r = 0.55)\). While 84% of subunits with rmsd values >2 Å and all subunits with rmsd values >5.1 Å have interfaces >2000 Å² (compared with 67% of all subunits), 62% of subunits with interfaces >2000 Å² have rmsds <2 Å. Thus, it appears that, while a large conformational change is generally associated with the formation of a large interface, a large interface alone is not necessarily a good indicator of conformational change.

One possible reason for the relatively weak correlation between interface size and rmsd is that this analysis only includes the conformational changes of individual subunits, whereas an interface is formed by two or more subunits. For a large interface to form, one can imagine that only a single binding partner would be required to undergo a large conformational change. However, this explanation does not apply to homomers due to their symmetric oligomerization. Since the correlation for homomers is only slightly greater than for heteromers, the contribution of this effect is likely small.

Relative Solvent Accessible Surface Area of Bound Subunits Is Predictive of Conformational Changes upon Binding

Given the limited correlation between interface size and rmsd, we sought to identify a more useful parameter for predicting conformational changes upon binding. Previous work has shown that the solvent accessible surface area \(A_s\) of a folded protein can be predicted with high accuracy from its molecular weight using a simple power-law relationship (Miller et al., 1987a, 1987b; Teller, 1976; Janin, 1976). In Figure S1 (available online), we plot molecular weight versus \(A_s\) for 4988 monomeric proteins. The fit to that plot results in Equation 1, where \(M\) is molecular weight. As expected, there is a high correlation between \(A_s\) and \(M\), and Equation 1 predicts \(A_s\) with an average absolute deviation of 5.8% for all monomers.
Protein Conformational Changes upon Binding

\[ A_s = 4.84M^{0.760} \]  

Since Equation 1 is highly predictive of the \( A_s \) of monomeric, folded proteins, we reasoned that the deviation of \( A_s \) from its predicted value could be a useful indicator of to what extent a protein resembles a folded monomer. Thus, we introduce the term relative solvent accessible surface area (\( A_{rel} \)) as the observed \( A_s \) scaled by its predicted value from Equation 1:

\[ A_{rel} = \frac{A_{observed}}{A_{predicted}} \]  

Importantly, we consider the \( A_{rel} \) value of each subunit in its bound conformation, but in isolation from the rest of the complex (\( A_{rel}(bound) \)). As an isolated subunit with a high \( A_{rel}(bound) \) value exposes more surface area than expected for a stable monomeric protein, we predict that significant conformational changes from its monomeric state are likely upon binding. This is similar to the previous observation that the subunits of some protein complexes would expose too much surface if their bound-state conformations were maintained in isolation, implying that intermolecular interactions must be responsible for their stabilization (Miller et al., 1987b).

In Figure 1B, we plot \( A_{rel}(bound) \) for each subunit from the monomer/complex data set versus the rmsd between bound and unbound conformations. The agreement is far better than for interface size, with correlations between \( A_{rel}(bound) \) and the observed rmsd of 0.82 for homomers and 0.83 for heteromers. If a linear relationship between \( A_{rel}(bound) \) and log(rmsd) is assumed, we obtain Equations 3 and 4.

\[ \text{RMSD}_{\text{homomer}} = \exp(6.14A_{rel}(bound) - 5.95) \]  
\[ \text{RMSD}_{\text{heteromer}} = \exp(6.35A_{rel}(bound) - 6.05) \]

We can also make more general statements about the relationship between \( A_{rel}(bound) \) and rmsd. For example, 79% of subunits with \( A_{rel}(bound) > 1.2 \) have rmsds > 5 Å (compared with 11% of all subunits) and 72% of subunits with \( A_{rel}(bound) > 1.1 \) have rmsds > 2 Å (compared with 30% of all subunits). Conversely, only 16% of subunits with \( A_{rel}(bound) < 1.1 \) and 6% of subunits with \( A_{rel}(bound) < 1 \) have rmsds > 2 Å. Thus, we suggest that \( A_{rel}(bound) \) values >1.2 can be used as strong indicators of large conformational changes upon binding (>5 Å rmsd), while subunits with \( A_{rel}(bound) < 1 \) are likely to undergo moderate conformational change (>2 Å rmsd). On the other hand, subunits with \( A_{rel}(bound) < 1.1 \) and especially <1 are much less likely to undergo significant conformational change.

It could be suggested that the correlation between \( A_{rel}(bound) \) and rmsd may arise from the presence of multidomain proteins in the data set. If these proteins have flexible linkers between more globular domains, this may cause these proteins to have larger \( A_{rel}(bound) \) values, while changes in the relative orientations of domains may lead to large rmsd values. We have addressed this in Figure S2 by repeating the analysis in Figure 1B using only proteins identified as having a single SCOP domain (Murzin et al., 1995). The correlations in this case are in fact even higher (0.88 for homomers, 0.84 for heteromers), demonstrating that the strong relationship between \( A_{rel}(bound) \) and rmsd is not due to flexible interdomain linkers.

Large Predicted Conformational Changes upon Binding Are Common

Given the apparent utility of \( A_{rel}(bound) \) for predicting protein conformational changes upon binding, we investigated the distribution of \( A_{rel}(bound) \) values in the full set of protein complexes from the PDB (after filtering, see Experimental Procedures). In Figure 2, we compare the distributions of \( A_{rel}(bound) \) values for the full set of complexes to those in the monomer/complex data set. For homomers, there is a slight, albeit not quite significant, increase in the average \( A_{rel}(bound) \) value, from 1.08 in the monomer/complex set to 1.10 in the full set of complexes (p = 0.07, Wilcoxon test). For heteromers, however, there is a very large increase in \( A_{rel}(bound) \) values, from 1.02 in the monomer/complex set to 1.14 in the full set (p = 2e-23).

The much larger \( A_{rel}(bound) \) values in the full set of heteromers compared with the monomer/complex data set suggest that binding-induced conformational changes are greater in general than might be expected from only considering complexes with corresponding monomeric structures available. For example, while only 6% of subunits from the monomer/heteromer set had \( A_{rel}(bound) \) values >1.2, more than four times as many (27%) have values that large in the full set of complexes. This suggests that large conformational changes upon binding are common, and that considering only those cases where a monomeric crystal structure is known may lead to bias against large
conformational changes in our understanding of molecular recognition mechanisms.

A simple explanation for the increased $A_{rel}^{(bound)}$ values in the full set of protein complexes comes from the hypothesis that the flexibility of unbound proteins correlates with the amount of conformational change they undergo upon binding. Since more flexible proteins are less likely to form diffraction-quality crystals, this would select against large conformational changes in the monomer/complex data set.

Intrinsically Disordered Proteins in Complex Have Larger $A_{rel}^{(bound)}$ Values Than Folded Proteins

Intrinsically disordered proteins are proteins that are completely disordered in their monomeric forms and which adopt highly extended conformations upon binding. Proteins predicted to be disordered (FoldIndex score of $s < 0$) and predicted to be folded (FoldIndex score of $s \geq 0$).

To determine to what extent intrinsic disorder contributes to the increased $A_{rel}^{(bound)}$ values observed in the full data set, we used the algorithm FoldIndex (Prilusky et al., 2005) to predict whether subunits were likely to be intrinsically disordered in isolation. FoldIndex provides a single score for an entire polypeptide chain, with values less than zero indicating that a protein is likely to be disordered. In Figures 3A and 3B, we split the subunits into two groups: predicted to be disordered (FoldIndex score $< 0$) and predicted to be folded (FoldIndex score $\geq 0$).

According to this classification, 4% of homomeric subunits and 13% of heteromeric subunits are disordered in isolation. Thus, if IDPs tend to have large $A_{rel}^{(bound)}$ values, this can explain the greater increase in $A_{rel}^{(bound)}$ values in the full data set versus the monomer/complex data set for heteromers compared with homomers.

The average $A_{rel}^{(bound)}$ values are significantly increased for both the disordered homomeric (1.24 versus 1.10, p = 3e-72, Wilcoxon test) and heteromeric (1.22 versus 1.13, p = 4e-79) subunits. Notably, there is a much greater fraction of proteins with very large $A_{rel}^{(bound)}$ values in the disordered subunits. We suggest that these high $A_{rel}^{(bound)}$ values correspond to proteins that are completely disordered in their monomeric forms and which adopt highly extended conformations upon

Differentiating between Structured and Flexible Binding

While $A_{rel}^{(bound)}$ is useful for predicting the rmsd upon binding for proteins with monomeric crystal structures available, in cases where the unbound state is undergoing large conformational fluctuations (i.e., proteins with high intrinsic flexibility or disorder), its utility is less clear. In other words, the concept of intrinsic disorder is often not applied to homomers, many homomeric complexes are known to undergo two-state folding in which the subunits are unfolded in isolation and only fold upon complex formation (Xu et al., 1998). MetJ is both predicted to be disordered (FoldIndex score of $s \approx -0.04$) and experimentally observed to be a two-state folder (Johnson et al., 1992). These observations are consistent with the fairly large $A_{rel}^{(bound)}$ value observed (1.22).

Figure 3D shows protein phosphatase 1 (PP1) in complex with the PP1-binding domain of spinophilin (PDB ID: 3EGG). This region of spinophilin is known to be highly disordered in isolation (Marsh et al., 2010; Ragusa et al., 2010) and is also predicted to be disordered (FoldIndex score of $s \approx -0.20$). It adopts a highly extended conformation in its complexed state with a very high $A_{rel}^{(bound)}$ value of 1.55, demonstrating the potential utility of this parameter for identifying intrinsically disordered regions.

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Figure 3. Intrinsically Disordered Proteins in Complex Have Large $A_{rel}^{(bound)}$ Values

(A) and (B) Distribution of $A_{rel}^{(bound)}$ values for (A) homomeric and (B) heteromeric subunits predicted to be intrinsically disordered and predicted to be bound.

(C) Homodimeric two-state folder (i.e., unfolded as a monomer) MetJ (A chain, red, B chain gray) (1CMB).

(D) PP1-binding domain of spinophilin which has been experimentally characterized as disordered in isolation (red), in complex with PP1 (gray) (3EGG).
A\(_{\text{ref}}\) of Free Proteins Correlates with Intrinsic Flexibility and Is Predictive of Conformational Changes upon Binding

\(A_{\text{ref}}\) values can only be calculated for complexes with 3D structures available. However, for many complexes, only structures of subunits in their monomeric states currently exist. In this section, we will assess the utility of \(A_{\text{ref}}\) for characterizing the intrinsic flexibility and conformational changes upon binding of monomeric proteins. Just as \(A_{\text{ref}(\text{bound})}\) values are useful for predicting the conformational changes of bound subunits, we propose that the \(A_{\text{ref}(\text{free})}\) values of free, monomeric proteins, may be useful for assessing the flexibility of their unbound states. That is, since proteins with higher \(A_{\text{ref}(\text{free})}\) values expose more surface area and adopt more extended conformations, they are likely to be more flexible.

In a previous study, Dobbins et al. performed normal mode calculations on a number of unbound proteins and compared them to their conformational changes upon binding (Dobbins et al., 2008). In Figure 5A, we compare the predicted flexibility values from their normal mode calculations against \(A_{\text{ref}(\text{free})}\) values for 60 monomers in their data set. We observe a significant correlation of 0.76, demonstrating that \(A_{\text{ref}(\text{free})}\) can indeed be used as a simple means of gauging intrinsic flexibility.

Given this correlation between \(A_{\text{ref}(\text{free})}\) and flexibility, we can now investigate the relationship between flexibility and binding-induced conformational changes in more detail. In Figure 5B, we have compared \(A_{\text{ref}(\text{free})}\) versus rmsd for all the...
monomer/heteromer pairs in our data set and observe a correlation of 0.80, which is only slightly less than was observed for \( A_{rel}(bound) \). Thus, the \( A_{rel}(free) \) values of unbound subunits appear to have significant utility for predicting conformational changes upon binding. In addition, this provides strong support for the idea that the magnitude of conformational changes upon binding correlates with intrinsic protein flexibility. Assuming a linear relationship between \( A_{rel}(free) \) and log(rmsd), we obtain Equation 5, which predicts all rmsds with an average absolute deviation of 37%.

\[
RMSD_{heteromer} = \exp(6.34A_{rel}(free) - 6.02) 
\]

As opposed to the strong correlation observed for heteromers, the correlation between \( A_{rel}(free) \) and rmsd for homomers is poor \((r = 0.18, \text{Figure S3})\). An explanation for this comes from the fact that most of the high \( A_{rel}(bound) \) subunits from the homomer data set are domain-swapped dimers (Liu and Eisenberg, 2002). In their domain-swapped conformations, these proteins extend large segments and tend to have high \( A_{rel}(bound) \) values, while in their monomeric forms, they resemble normally folded proteins with low \( A_{rel}(free) \) values. Thus, the \( A_{rel}(free) \) values of monomeric proteins are not useful for predicting the conformational changes they undergo upon self-association.

In Figure S4, we have compared both predicted flexibility values from normal mode calculations and \( A_{rel}(free) \) values to rmsd values for the 60 monomer/complex pairs from the study by Dobbins et al. (all complexes from this study were heteromers) (Dobbins et al., 2008). Predicted flexibility values have a correlation of 0.43 with rmsd, confirming the association between flexibility and conformational change previously observed (Dobbins et al., 2008). Notably, the correlation for \( A_{rel}(free) \) is even higher \((r = 0.55)\). These results strongly support the use of \( A_{rel}(free) \) as a simple proxy for the flexibility of free proteins and for predicting the magnitude of binding-induced conformational changes, with \( A_{rel}(free) \) being very simple to compute and providing information complementary to normal mode analysis.

One possible way of viewing \( A_{rel} \) may be that it is, to a certain extent, acting as a measure of protein globularity. Thus, a simple way of interpreting our results is that more globular proteins tend to have less solvent accessible surface area, are less flexible while free in solution, and undergo smaller conformational changes upon binding. Note that this is independent of whether the protein is single or multidomain.

**Examples of Proteins Undergoing Conformational Change**

In Figure 6, we show three examples of proteins in their bound and unbound states in order to illustrate the relationships between \( A_{rel}(bound) \), \( A_{rel}(free) \) and conformational changes upon binding. Figure 6A shows the 16S ribosomal RNA processing protein RimM in its free state (PDB ID: 2DYI) and in complex with the ribosomal protein S19 (PDB ID: 3A1P). Bound RimM has an \( A_{rel}(bound) \) value of 1.16, consistent with the conformational change observed upon binding (rmsd of 3.9 Å), which primarily involves rearrangement of loops and β strands in the one domain that directly interacts with S19. The \( A_{rel}(free) \) value of 1.16 indicates significant free-state flexibility that is likely important for facilitating binding.

Figure 6B shows calmodulin (PDB ID: 4CLN) and its complex with myosin VI (PDB ID: 2VB6). Here, we see a large \( A_{rel}(bound) \) value (1.27), consistent with the major conformational change upon binding (rmsd of 15.0 Å). Furthermore, calmodulin also has a very large \( A_{rel}(free) \) value (1.30), due to its highly extended monomeric structure, implying considerable intrinsic flexibility. Indeed, this is supported by the large number of structures of varying conformations available for both free and bound states of calmodulin, which is likely due to this high flexibility (Chou et al., 2001; Gsponer et al., 2008).

Figure 6C shows thioredoxin in both monomeric (PDB ID: 2O7K) and domain-swapped dimeric states (PDB ID: 3DIE). The \( A_{rel}(bound) \) of bound thioredoxin is very large (1.43) and so is the conformational change between monomeric and dimeric states (rmsd = 15.0 Å). However, \( A_{rel}(free) \) is quite low (0.88), implying a highly stable monomeric state. This is consistent with the poor correlation between \( A_{rel}(free) \) and rmsd mentioned...
earlier for homomers. In this case, the domain-swapped dimer of thioredoxin only forms upon mutation. Thus, as is probably true for most proteins in the monomer/homomer set, sequence differences and/or differences in sample conditions, and not intrinsic flexibility, account for the large conformational differences between free and bound states.

As an interesting note, during the course of our study, we encountered a pair of structures (monomeric 1CMW and hetero-dimeric 1BGX) where an $A_{rel}^{\text{bound}}$ of 1.38 and an $A_{rel}^{\text{free}}$ of 1.29 predict a very large conformational change upon binding. However, the free and bound states adopted very similar conformations (rmsd of only 1.4 Å) and examination of Figures 1B and 3A shows us that there are no other proteins near these regions of the plots. Further investigation revealed that the monomeric structure 1CMW had been recently retracted from the PDB and is believed to have been fabricated (presumably using the complex structure as a template). Thus, in this case, $A_{rel}$ clearly proved its usefulness by identifying an implausible structure.

**Conclusion**

In this study, we have presented simple methods for predicting the binding-induced conformational changes of proteins using the relative solvent accessible surface area of either complexed or monomeric structures. Although $A_{rel}^{\text{bound}}$ and $A_{rel}^{\text{free}}$ provide no information on the nature of the conformational change, only a predicted magnitude, this will allow expanded data sets and facilitate large-scale analyses of the relationships between protein sequences, structures, flexibility and conformational changes upon binding.

The use of $A_{rel}^{\text{bound}}$ to predict conformational changes upon binding will be particularly useful for the large number of protein complexes for which no structures of the free subunits have been determined. Thus, one can quickly and easily assess whether an unbound subunit is likely to resemble its bound conformation or whether it would require significant conformational changes. Furthermore, one can consider the likelihood that a bound subunit is highly flexible in its monomeric state: an $A_{rel}^{\text{bound}}$ value $>$1.2 indicates a high chance of significant flexibility, while $A_{rel}^{\text{bound}}$ $>$1.4 suggests a protein is very likely to be intrinsically disordered. $A_{rel}^{\text{bound}}$ could also have considerable utility as a structure evaluation tool. For example, in a docking calculation, the predicted complex structure could be compared with the unbound structure to assess whether the amount of conformational change is reasonable. $A_{rel}^{\text{free}}$ can be used to predict both the flexibility of free proteins and their conformational changes upon binding. This could also be very useful for the docking of protein complexes: since complexes that require large conformational changes from the monomeric state are much less likely to be successfully modeled, this could allow the probability of a successful docking calculation to be predicted beforehand. In addition, knowledge of the expected conformational change from $A_{rel}^{\text{free}}$ could also potentially be useful for guiding docking calculations.

There is growing evidence that many proteins can undergo motions over a wide range of length and timescales and that these motions are often important for biological function. Similarly, along with the large body of recent research into the importance of protein disorder has come the awareness of the continuum of protein flexibility and disorder, with some disordered states possessing large amounts of nonrandom secondary and tertiary structure and some folded proteins being highly flexible (Boehr et al., 2009; Gsponer et al., 2008; Jensen et al., 2009; Marsh et al., 2010). Our results support this paradigm by showing that proteins undergo widely varying amounts of conformational change upon binding, and by suggesting that the magnitude of this conformational change tends to correlate with flexibility and intrinsic disorder in the unbound state. Furthermore, a significant correlation between protein flexibility and conformational changes upon binding supports the importance of conformational selection in molecular recognition. The observation that flexible proteins tend to undergo greater conformational changes upon binding suggests that this intrinsic flexibility is important for sampling those conformations resembling the bound state.

Some complexes that lack corresponding monomeric structures are probably obligate (Nooren and Thornton, 2003), i.e., unstable on their own, and thus rarely or never exist in a monomeric state within the cell. Indeed, this could be true for a considerable fraction of the subunits we predicted to undergo flexible binding. For these cases, it might be argued that the ability to predict the conformational changes required of a folded monomeric state would be of limited utility. However, even if a complex is completely obligatory, at some point after polypeptide synthesis the complex must initially assemble and these conformational changes must be important for the assembly process. This does raise the question, however, of how to interpret $A_{rel}^{\text{bound}}$ values for proteins which undergo significant changes in flexibility upon binding. Can the predicted rmsd values be compared with proteins that undergo structured binding? To what extent do $A_{rel}^{\text{bound}}$ values correlate with changes in structure and dynamics for flexible proteins?

Since our predictive method is based upon only complexes which have paired monomer crystal structures, it is difficult to assess these questions at this point. Moreover, while the $A_{rel}$ method is currently only applicable to crystallizable protein complexes, recent work has demonstrated that some proteins can remain highly flexible or disordered in their complexed states (Fong et al., 2009; Mittag et al., 2010a; Tompa and Fuxreiter, 2008). Such dynamic complexes are largely inaccessible to X-ray crystallography, although they are now in some cases able to be characterized in significant detail by small-angle X-ray scattering and NMR (Marsh et al., 2010; Mittag et al., 2010b; Wells et al., 2008). Future advances in computational and experimental modeling of highly flexible proteins, and especially the development of methods that can be applied on a large scale, will be crucial for fully understanding the range of protein interactions and conformational dynamics utilized within the cell.

**EXPERIMENTAL PROCEDURES**

We selected all structures from the full set of protein X-ray crystal structures in the PDB with PISA (Krissinel and Henrick, 2007) quaternary structure assignments determined to a resolution $\leq 2.5$ Å (2010-11-14). Only monomers and subunits longer than 40 residues were considered. Monomers with more than five disordered residues (defined as nonterminal residues not observed in the crystal structure) were ignored. In addition, we filtered out all monomers with $\geq 25$ nonwater HETATM lines in order to remove most monomers with ligands or significant chemical modifications. To ensure that our data set contained only high-confidence monomers, we only considered monomers...
identified by PISA where both the asymmetric unit and the PDB biological unit contained single chains, and which were not identified as having more than one subunit in the PIQSi database of manually curated quaternary structure (Levy, 2007). We also filtered out protein complexes that had only a single chain in either their asymmetric unit or PDB biological unit, or that were identified as monomers in PIQSi. Complexes containing only a single polypeptide chain longer than 20 residues were removed in order to filter out single protein chains bound to small peptides or nucleic acids. Finally, we removed complexes that were formed by polypeptide cleavage. These were identified as complexes containing unique chains that shared the same UniProt identifiers but which had <90% sequence identity to each other. Together, this filtered set of protein complexes comprised the “full set” referred to in the text.

To generate the monomer/complex pairs data set, a protein BLAST (Altschul et al., 1997) search was performed for all monomers against all unique subunits in multimeric complexes. Only pairs with >98% sequence identity and similar rmsd values (<1 Å difference) were dropped. The solvent accessible surface area of each structure and isolated subunit was calculated with AREAIMOL (Collaborative Computational Project, 1994). The interface size of each subunit was calculated as the amount of surface area buried between the subunit and the rest of the complex. Complexes were identified as homomeric if all of the protein subunits had ≥90% sequence identity to each other; otherwise they were heteromeric. All pairs from the monomer/complex data set along with their $A_{\text{binding}}$, $A_{\text{free}}$, interface size and rmsd values are provided in Tables S1 and S2.

### SUPPLEMENTAL INFORMATION

Supplemental information includes four figures and two tables and can be found with this article online at doi:10.1016/j.str.2011.03.010.

### ACKNOWLEDGMENTS

We thank Emmanuel Levy for helpful comments on the manuscript. J.A.M. was supported by a European Molecular Biology Organization Long-Term Fellowship.

Received: February 3, 2011
Revised: March 14, 2011
Accepted: March 22, 2011
Published: June 7, 2011

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