

# A structure-based benchmark for protein–protein binding affinity

Panagiotis L. Kastritis,<sup>1</sup> Iain H. Moal,<sup>2</sup> Howook Hwang,<sup>3</sup> Zhiping Weng,<sup>3</sup> Paul A. Bates,<sup>2</sup> Alexandre M. J. J. Bonvin,<sup>1</sup> and Joël Janin<sup>4\*</sup>

<sup>1</sup>Bijvoet Center for Biomolecular Research, Faculty of Science, Utrecht University, 3584CH Utrecht, The Netherlands

<sup>2</sup>Biomolecular Modelling Laboratory, Cancer Research UK London Research Institute, Lincoln's Inn Fields Laboratories, London WC2A 3LY, United Kingdom

<sup>3</sup>Program in Bioinformatics and Integrative Biology, University of Massachusetts Medical School, Worcester, Massachusetts 01605

<sup>4</sup>Yeast Structural Genomics, IBMC UMR 8619, Université Paris-Sud, 91405 Orsay, France

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**Abstract:** We have assembled a nonredundant set of 144 protein–protein complexes that have high-resolution structures available for both the complexes and their unbound components, and for which dissociation constants have been measured by biophysical methods. The set is diverse in terms of the biological functions it represents, with complexes that involve G-proteins and receptor extracellular domains, as well as antigen/antibody, enzyme/inhibitor, and enzyme/substrate complexes. It is also diverse in terms of the partners' affinity for each other, with  $K_d$  ranging between  $10^{-5}$  and  $10^{-14}M$ . Nine pairs of entries represent closely related complexes that have a similar structure, but a very different affinity, each pair comprising a cognate and a noncognate assembly. The unbound structures of the component proteins being available, conformation changes can be assessed. They are significant in most of the complexes, and large movements or disorder-to-order transitions are frequently observed. The set may be used to benchmark biophysical models aiming to relate affinity to structure in protein–protein interactions, taking into account the reactants and the conformation changes that accompany the association reaction, instead of just the final product.

**Keywords:** protein–protein interaction; molecular recognition; binding free energy; conformation changes; allostery

## Introduction

Relating structure to function is a major objective of structural biology. The noncovalent interactions

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Additional Supporting Information may be found in the online version of this article.

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\*Correspondence to: Joël Janin, Joël Janin, IBMC Bat. 430 Université Paris-Sud, 91405-Orsay, France.  
E-mail: joel.janin@u-psud.fr

many proteins make with other biomolecules are essential to their function. Protein–protein interactions build assemblies that are as diverse as life itself.<sup>1–3</sup> When a high-resolution structure is available, the interaction can be described in atomic details, but it is the affinity of the components for each other that determines whether the assembly actually exists under a given condition of temperature, pH, and protein concentration, and whether it is transient or permanent. For a binary complex, the binding affinity translates in physical–chemical terms into an equilibrium dissociation constant ( $K_d$ ), which may be measured at equilibrium or derived from the reaction kinetics, and the related Gibbs free energy of dissociation  $\Delta G$ .  $K_d$  measurements have been performed on many protein–protein

complexes that also have a X-ray structure, and several models have attempted to correlate affinity to features observed in these structures. Although some have been very successful on small training sets,<sup>4,5</sup> the published models did far less well on larger sets,<sup>6–10</sup> and their predictive value remains poor.<sup>11</sup>

The bearing of such studies depends on the quality as well as the size of the set of experimental data on which the models are tested, and on the diversity of the biological systems they represent. Whereas early structural data concerned almost exclusively protease/inhibitor and antigen/antibody complexes, many more structures are now available, and the recent sets are more diverse. However, associating a crystal structure to biophysical measurements done in solution is an error-prone process, and the published sets contain many incorrect affinity data.<sup>11</sup> Moreover, the structural data in these sets represent the complexes (the “bound” structures), but not their free components (the “unbound” structures). Therefore, the models based on them describe the thermodynamics of association reaction by its product only, ignoring the reactants and the structure changes they may undergo.

A benchmark set of validated  $K_d$  and  $\Delta G$  data that can be reliably assigned to the bound and unbound structures of a complex should be of great value when modeling recognition. The one we present here is the product of collaboration between four research groups, checking each other's data. It contains 144 entries that represent many different biological systems with affinities that cover nine orders of magnitude in  $K_d$ , and structures that display a wide variety of conformation changes.

## Results

### The data

The starting point of the present study was the docking benchmark Version 4.0 (DB4.0), which contains protein data bank (PDB<sup>12</sup>) entries for 176 protein–protein complexes and their unbound components.<sup>13</sup> The complexes are nonredundant, they have a X-ray structure solved at better than 3.25 Å resolution, and with the exception of a few antibodies, their unbound components have either a X-ray or a NMR structure.

We found affinity data in the literature for 144 protein–protein complexes; 130 are from DB4.0, and the rest is closely related to entries in it; seven replace entries for which we could not find a  $K_d$ , and seven are new. The experimental methods used to determine the  $K_d$  are highly dependent on the proteins and on their affinities. Surface plasmon resonance (SPR), isothermal calorimetry (ITC), and titration by fluorescence or other spectroscopic methods are all applicable if the  $K_d$  is in the micromolar to nanomolar

range. Together, they yielded 75% of the data we retained; enzymatic inhibition studies yielded 14%.  $K_d$  values are usually reported in publications with standard errors of 20–50%, equivalent to 0.1–0.25 kcal mol<sup>-1</sup> for  $\Delta G$ . When  $K_d$  is below nanomolar, ITC is limited by its sensitivity, and SPR by the slow reaction kinetics that make the measurements less reliable. Moreover, each method has specific drawbacks and requirements, such as a reporter group in fluorimetry, protein immobilization in SPR, or the kinetic model of the reaction needed to convert an inhibition constant ( $K_i$ ) into a  $K_d$ . About half of the entries in our set have corroborating values determined by two or more methods under similar conditions. They commonly differ by a factor of 2 for  $K_d$ , or 0.4 kcal mol<sup>-1</sup> for  $\Delta G$ , which is a more realistic estimate of the standard error than that obtained with a single method.

The discrepancy can be much greater when the experimental conditions differ. The dependence of  $K_d$  on temperature, ionic strength, and pH has been checked in a number of cases, which we will assume to be representative of the whole set. All the measurements we report except three have been done in the temperature range 18–35°C. The data on the dissociation enthalpy  $\Delta H$ , determined either by ITC or the van't Hoff equation, suggest that the correction does not exceed a factor of 2 in this range (e.g.,  $|\Delta H| < 10$  kcal mol<sup>-1</sup>). Changing the ionic strength in the range 0.1–0.5M can have a larger effect, but in general, pH is the most significant environmental factor. If ligand binding induces a pK shift in an acid or base group that has a pK near the pH of the experiment, that group will take or release protons as the complex forms, and thus,  $K_d$  will depend on the proton concentration. Changing the pH in the range 5.5–8.5, which covers 96% of our data, can change  $K_d$  by a factor of 10–50, and  $\Delta G$  by 1.4–2.3 kcal mol<sup>-1</sup>, which largely exceeds any effect of temperature or ionic strength. Moreover, the dependence of  $K_d$  on pH is just one example of allosteric effects discussed below, by which the concentration of one ligand, proton in this case, changes the affinity of a protein for another ligand.

### Classes of complexes and affinities

Of the 144 protein–protein complexes of our set, all but 7 fulfill the condition initially set for the docking benchmark, that the PDB should contain entries for both the assembly and its unbound components. The exceptions are antigen/antibody complexes that were included in DB4.0 even though the antibody has no unbound structure. In Table I, the set is broken into the same three functional classes as in the docking benchmark: (A) antibody/antigen, (E) enzyme containing, and (O) other complexes; subclasses are introduced in the latter two. In addition, the set is split into three categories based on  $K_d$ : high

**Table I.** *Classes of Complexes*

Class	Number		Affinity class <sup>a</sup>			$\Delta G$ (kcal mol <sup>-1</sup> ) <sup>b</sup>		Large conformation changes <sup>c</sup>
	All	Noncognate	High	Medium	Low	Mean	S.D.	
A								
Antigen-antibody	19	2	2	16	1	12.2	1.3	0
E								
Enzyme/inhibitor	40	4	17	22	1	13.8	2.3	7
Other enzyme complexes	21	1	0	12	9	9.2	1.9	7
O								
G-proteins	17	—	1	6	10	8.9	2.5	6
Receptors	13	—	1	11	1	11.5	2.1	4
Miscellaneous	34	2	0	22	12	9.3	2.2	11
All	144	9	20	90	34	11.0	2.9	35

<sup>a</sup> High  $K_d < 10^{-10}M$ , medium  $10^{-6}$  to  $10^{-10}M$ , and low  $K_d > 10^{-6}M$ .

<sup>b</sup> Mean value and standard deviation excluding the noncognate complexes.

<sup>c</sup> Complexes with  $I_{\text{rmsd}} > 1.5 \text{ \AA}$ .

( $K_d < 0.1 \text{ nM}$ ), medium (0.1 nM to 1  $\mu M$ ), and low affinity ( $>1 \mu M$ ). They represent 14%, 63%, and 23%, respectively, of the cases. In addition, Table I mentions the presence of 9 noncognate complexes (see below) and provides the mean values of  $\Delta G$  and their standard deviations in each class, calculated on the cognate complexes only.

Class A contains monoclonal antibodies, fairly similar in terms of their affinity for the protein antigens they were raised against. All but three of the 19 class A complexes are in the medium-affinity category; their  $K_d$ 's are in the range 0.1–70 nM. The exceptions are the noncognate complex 2VIS (see below), which is of low-affinity, and two complexes with a high affinity. Fab BO2C11, which has a picomolar  $K_d$  for Factor VIII (1IQD), was not obtained by clonal selection in hybridoma cell cultures like the other antibodies in our sample; it is the product of a cell line derived from the memory B-cell repertoire of a patient with hemophilia A.<sup>14</sup>

Class E, enzyme-containing complexes, is highly heterogeneous in comparison to class A. However, when the complexes with inhibitors are set apart from those with substrates or regulatory subunits (“other enzyme complexes” in Table I), it becomes obvious that the former have a much higher affinity. Indeed, all but four of the high-affinity complexes are enzyme/inhibitor. Five have a  $K_d$  below 0.1 pM, at the lower limit of what can be reliably measured given the very low dissociation rate that such a  $K_d$  implies. These complexes, stable on a time scale of days or even months, implicate the protease trypsin, three nucleases, and uracyl-DNA glycosylase.

Class O (“other”) was introduced in the docking benchmark for convenience at a time where most of the available structural data concerned antigen/antibody and enzyme/inhibitor complexes.<sup>15</sup> It has greatly expanded since then, and now comprises 64 complexes that perform all sorts of biological functions. Among them, 17 form a subclass that contains G-proteins. These proteins bind GTP and hydrolyze

it to GDP; they play a central role in signal transduction, membrane traffic, and other cellular processes. They have many partners: GTPase activating proteins (GAPs) that enhance their GTPase activity, guanine nucleotide exchange factors that allow the GDP product to be released, protein kinases, etc. Only one of the complexes of this subclass is high-affinity (1I2M, Ran/RCC1); the others have  $K_d$  in the nanomolar to low micromolar range, consistent with functions that require the interaction to be transient on a time scale of seconds to minutes. Another subclass, with 13 members, contains cell surface receptors, present mostly as the extra cellular domain binding a cytokine or protein hormone. All but two are of medium-affinity. The 34 “miscellaneous” complexes that remain carry highly diverse functions: 12 are low-affinity, and two have  $K_d > 0.1 \text{ mM}$ , at the upper limit of what can be reliably measured. These very low affinity complexes implicate ubiquitin (1S1Q) and the T-cell receptor (1AKJ). Two other low affinity complexes (1XD3, 2OOB) contain ubiquitin.

### **Cognate and noncognate protein–protein association**

Table II lists nine pairs of entries representing closely related proteins that form complexes of a similar geometry, but with a very different  $K_d$ . The more affine in each pair is labeled as cognate, because in most cases, it is the one of biological relevance. Thus, the cognate partner of the *Bacillus amyloliquefaciens* inhibitor barstar, is the nuclease barnase produced by that bacterium, for which it has femtomolar affinity<sup>26</sup>; barstar also inhibits RNase SA from *Streptomyces aureofaciens*, but with nanomolar affinity only. The colicin/immunity protein system provides some remarkable examples of cognate vs. noncognate assemblies,<sup>27</sup> one of which is cited in Table II and illustrated by Figure 1. The strain of *Escherichia coli* that makes colicin E9, endowed with a DNase activity, also produces the Im9 immunity protein that inhibits it very

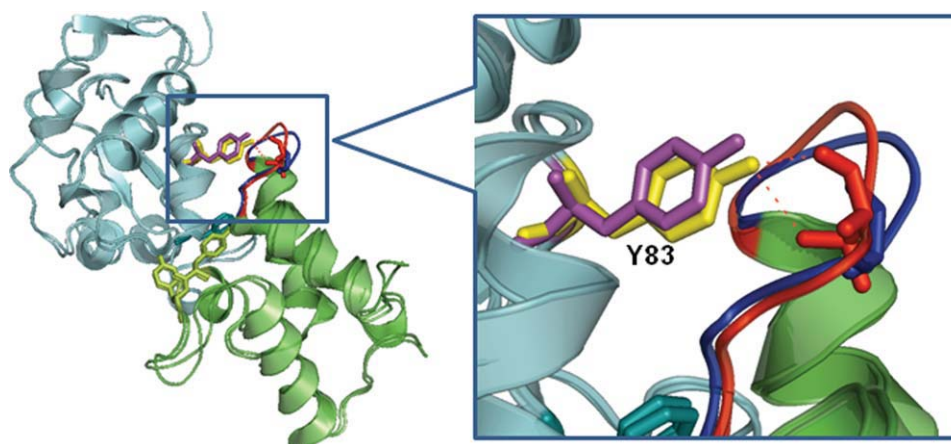
**Table II.** Cognate vs. Noncognate Complexes

Cognate			Noncognate			$\Delta\Delta G$ (kcal mol <sup>-1</sup> )	References
PDB entry		$K_d$ (M)	PDB entry		$K_d$ (M)		
Enzyme/inhibitor							
1BRS	Barnase/barstar	5.0E-14	1AY7	RNase SA/barstar	1.0E-10	4.1	16
1EMV	Colicin E9/Im9	2.4E-14	2WPT	Colicin E9/Im2	1.5E-8	7.9	17
2PTC	Trypsin/BPTI	6.0E-14	1CBW	Chymotrypsin/BPTI	9.0E-9	7.3	18
2PTC	Trypsin/BPTI	6.0E-14	2TGP	Trypsinogen/BPTI	2.3E-6	10.5	19
Enzyme/substrate							
2PCC	Yeast peroxidase/ yeast cyt <i>c</i>	1.6E-6	2PCB	Yeast peroxidase/ horse cyt <i>c</i>	1.0E-5	1.1	20,21
Antigen/antibody							
2VIR	Flu hemagglutinin/ Fab HC19	1.0E-9	2VIS	Flu HA (T131I)/Fab HC19	4.0E-6	4.9	22
1P2C	Fab F10.6.6	1.0E-10	1MLC	Fab D44.1/lysozyme	7E-8	3.9	23
Miscellaneous							
1EFN	HIV-1 Nef/Fyn SH3 (R96I)	3.8E-8	1AVZ	HIV-1 Nef/wild type Fyn SH3	1.6E-5	3.6	24
3BZD	TCR-V $\beta$ /SEC3-1A4 variant	9.6E-8	2AQ3	TCR-V $\beta$ /wt SEC3	1.2E-5	2.9	25

efficiently; a different strain produces the Im2 immunity protein, which has a much lower affinity for E9.<sup>29</sup> Like the immunity proteins, the bovine pancreatic trypsin inhibitor (BPTI) has a femtomolar  $K_d$  for the cognate serine protease trypsin. BPTI also binds trypsinogen, the inactive precursor of trypsin,<sup>30</sup> and it inhibits chymotrypsin, a related protease with a different specificity, but its affinity for these two proteins is five to eight orders of magnitude less than for trypsin. In all four pairs of enzyme/inhibitor complexes, sequence changes introduce several residue substitutions at the interface. In cytochrome peroxidase/cytochrome *c*, an enzyme/substrate complex,  $K_d$  changes by a factor of 6 only,

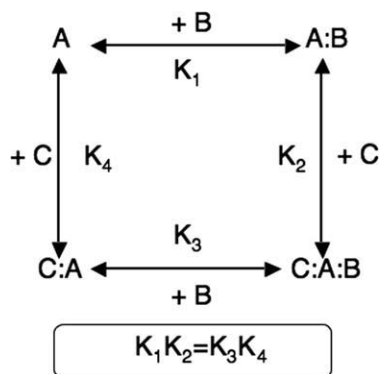
when horse cytochrome *c* replaces the cognate yeast substrate, even though the sequence identity is only 35%. Perhaps more significantly, the stoichiometry of the crystalline complex changes from 1:1 to 2:1.<sup>31</sup>

Table II also cites two systems in which a single residue substitution leads to a large change in  $K_d$ . Antibody HC19, raised against flu hemagglutinin, has a high affinity for the wild type protein, and a much lower one for a point mutant that allows the virus to escape neutralization by the antibody in cell cultures.<sup>22</sup> Protein Nef of HIV-1 forms a complex with both the SH3 domain of Fyn, a tyrosine kinase that binds Nef poorly, and a variant in which a point substitution has been introduced to mimic Hck,



**Figure 1.** Cognate and noncognate colicin/immunity protein complexes. The DNase domain of colicin E9 (cyan) has a very high affinity for the Im9 immunity protein produced by the same *E. coli* strain (green), and a 10<sup>6</sup>-fold lower affinity for Im2, produced by a different strain.<sup>17</sup> The crystal structures of the cognate E9/Im9 (1EMV) and the noncognate E9/Im2 (2WPT) complexes indicate that the mode of assembly is essentially the same. The insert shows that, nevertheless, segment 22–30 of Im9 (red) interacting with Tyr83 of E9 undergoes a significant movement in Im2 (blue), where it has a different sequence.<sup>28</sup>





	2TPI	1H9D	1K5D
A	Trypsinogen	CBFa (Runx1)	Ran.GTP
B	BPTI	CBFβ	RanGAP
C	IleVal	DNA	RanBP1
$K_1$	2.3E-6	<b>5.2E-7</b>	7E-6
$K_2$	1E-5	1.9E-12	<b>3E-10</b>
$K_3$	<i>6E-14</i>	<b>4.5E-8</b>	2E-6
$K_4$	<b>400</b>	2.3E-11	1E-9
Ratio	4E7	11	3.5

**Figure 2.** Allosterity and ligand affinity. Protein A binds ligands B and C at two distinct sites that have different affinities in the binary and the ternary complexes. The ratio  $K_1/K_3 = K_4/K_2$  is a measure of the cooperative interaction between the ligands. The table reports dissociation constants (in molar units); they are from experiment, except for  $K_3$  in 2TPI, which is an estimate based on trypsin, and the three values in bold face, calculated from the linkage equation; the values retained for the benchmark are underlined. References: 2TPI,<sup>19</sup> 1H9D,<sup>47,48</sup> and 1K5D.<sup>49</sup>

which binds Nef much more tightly than Fyn.<sup>32</sup> Here again, the stoichiometry in the crystal changes from 1:1 to 2:1 between the higher and the lower affinity complex. The mutation in Fyn was introduced by site-directed mutagenesis, which has been a standard method to study recognition and specificity for more than 20 years.<sup>33,34</sup> Most of the substitutions made at a protein–protein interface in this way cause a loss of affinity, locating “hot spots” at the interface when the effect is large.<sup>35</sup> However, a change in affinity can also point to a change in conformation, which can be assessed if the mutant structure has been determined in both the unbound and the bound state. This has been done for a few systems that are represented in this benchmark by the wild-type only: barnase/barstar, TEM1-BLIP, lysozyme/antibody, complement C3/Efb-C.<sup>36–39</sup> For the mutants, we refer the reader to the publications, and to databases such as PINT, ASEdb, and PDBbind.<sup>40–42</sup>

Mutations that improve affinity as in Fyn SH3 are much less common, and therefore of particular interest. They play an essential part in the maturation of antibodies by allowing the affinity for the antigen to increase beyond the initial clonal selection step.<sup>43</sup> The high affinity of antibody BO2C11 for Factor VIII mentioned above can be explained in this way.<sup>14</sup> The effect of maturation on affinity has been analyzed in many systems, and its effect on structure, in several antibody/lysozyme complexes.<sup>23,44,45</sup> One is cited in Table II: the low-affinity D44.1 and the high-affinity F10.6.6 monoclonal antibodies derive from the same germ-line genes, but F10.6.6 was raised after long-term exposure to the antigen.<sup>23</sup> Table II contains another pair of complexes that is relevant to the immune system. When the superantigen SEC3 (*Streptococcus* exotoxin C3) binds to the Vβ chain of the T-cell receptor, the wild type protein has  $K_d > 1 \mu M$ ; the 1A4 variant and other variants selected by phage display have a much better affinity.<sup>25</sup>

### Allosterity

A number of the proteins in the benchmark are allosteric in the original sense of the term.<sup>46</sup> They carry other binding sites than the one for which we report a  $K_d$  and have several partners: metal ions, small molecules, nucleic acids or proteins. If a conformation change accompanies the binding of one ligand, the affinity for other ligands may be greatly affected.

Trypsinogen is a remarkable example. This precursor of trypsin is catalytically inactive, yet it binds BPTI in the same way as trypsin, albeit with a  $K_d$  that is orders of magnitude higher<sup>19</sup> (Table II). The structures of the trypsin/BPTI and trypsinogen/BPTI complexes are very similar,<sup>30</sup> and they offer no explanation for the change in affinity. That of unbound trypsinogen does: the inactive precursor contains disordered loops that become ordered when it is activated to trypsin, and also when it binds BPTI. The activation involves the proteolytic cleavage of a peptide bond. This releases a free amino group that can interact with the aspartate residue adjacent to the catalytic serine on the opposite face of the protein, triggering the conformation change. Adding the IleVal dipeptide, which mimics the N-terminal sequence of active trypsin, allows the same interaction to occur, and greatly increases the trypsinogen affinity for BPTI. The trypsinogen/IleVal/BPTI ternary complex has again the same structure as trypsin/BPTI.<sup>19</sup> In the reaction scheme of Figure 2, inspired from,<sup>19</sup> only  $K_1$  and  $K_2$  have effectively been measured. The value of  $K_3$  assumes that trypsinogen/IleVal has the same affinity as trypsin for BPTI, and  $K_4$  is derived from it through the linkage equation. If trypsinogen/IleVal has less affinity for BPTI than trypsin, which is likely,  $K_4$  may be less than calculated, but it is too high in any case to be measured or allow an X-ray structure to be determined for the binary trypsinogen/IleVal complex. As a consequence, the ternary complex cannot be included in the benchmark.

A few ternary complexes are nevertheless present. In the complex of the Ras G-protein with the Son of Sevenless exchange factor (1NVU), two Ras molecules occupy two noninteracting sites on the exchange factor.  $K_d$ 's being available for each site,<sup>50</sup> we made two entries for this complex. Two other ternary complexes are cited in Figure 2. Their sites do interact, albeit not as strongly as in trypsinogen. One (1H9D) contains the  $\alpha$  subunit of the core-binding factor (CBF $\alpha$ ), the  $\beta$  subunit (CBF $\beta$ ), and DNA.<sup>51</sup> The affinity of CBF $\alpha$  for DNA has been measured both in the absence of CBF $\beta$  and in its presence, where it goes up 11-fold. The  $K_d$  reported in the benchmark is  $K_3$ , measured by titrating CBF $\alpha$ /DNA with CBF $\beta$ <sup>47</sup>; the linkage equation predicts that  $K_1$  must be less by the same factor in the absence of DNA. The other ternary complex (1K5D) involves the Ras related G-protein Ran, bound to RanGAP and to RanBP1.<sup>52</sup> Seewald *et al.*<sup>49</sup> report  $K_d$ 's for RanGAP in complex with Ran and with Ran/RanBP1; they differ by a factor of 3.5. The  $K_d$  in the benchmark is  $K_2$ , calculated by dividing the experimental value of  $K_4$  by that factor. This system contains another very important component: the nucleotide. It is a GTP analog in these Ran structures and in the  $K_d$  measurements reported in Table II.  $K_4$  changes by four orders of magnitude if GDP replaces GTP.<sup>49</sup> In the high-affinity complex with RCC1 (1I2M), Ran contains no nucleotide. The Ran affinity for RCC1 is much less when it binds GDP, and the difference contributes to the nucleotide exchange mechanism.

In all G-proteins, the presence and nature (GDP or GTP) of the nucleotide affects both the protein conformation and its affinity for its partners. Metal ions, especially  $\text{Ca}^{++}$ , play a similar role in other systems. We therefore checked that the  $K_d$ 's have been measured in the presence (or absence) of the same nucleotide or metal ion, as the X-ray structures.

### Conformation changes

Ligand binding can affect the protein conformation in many ways. Side chain rotations and small local adjustments of the main chain always take place, and large movements of surface loops are common. On a more global scale, whole domains or whole subunits may rotate or shift, and part of the polypeptide chain may change secondary structure or undergo a disorder-to-order transition.<sup>53,54</sup> Examples of all those are present in the benchmark, and can be identified by comparing the unbound proteins to the complexes.

In line with the docking benchmark,<sup>13</sup> we use the  $I_{\text{rmsd}}$  (interface C $\alpha$  root-mean-square displacement) parameter to detect conformation changes and estimate their amplitude. In our set, only 26 (18%) of the complexes have  $I_{\text{rmsd}} < 0.5 \text{ \AA}$ , which war-

rants that the main chain has essentially the same conformation bound and unbound, and thus, that the components associate as rigid bodies; two-thirds of those are antigen/antibody or enzyme/inhibitor complexes. In all the antigen/antibody complexes, the antigen moiety undergoes changes of limited amplitude only; the antibody does too, at least in the 12 complexes where its unbound structure is known.

A majority of the complexes (83%, or 58%) have  $I_{\text{rmsd}}$  values in the range 0.5–1.5  $\text{Å}$ , meaning that significant movements take place at the interface, but no large scale ones. Then, 35 complexes (24%) have  $I_{\text{rmsd}} > 1.5 \text{ \AA}$  (Table I), and they display major conformation changes. Whereas many protein inhibitors bind the enzyme as a rigid body, several enzyme/inhibitor complexes undergo large changes. In trypsinogen/BPTI (2PTC), BPTI is rigid, but trypsinogen undergoes a disorder-to-order transition that affects several surface loops.<sup>55</sup> In four other systems (1JIW, 1ZLI, 2O3B, and 4CPA), the inhibitor interacts with the enzyme through N- or C-terminal segments that are disordered in NMR structures of the unbound protein, and become ordered in the complex. In the ClpA/ClpS complex (1R6Q), the N-terminal segment of ClpS makes up much of the contact with ClpA, yet it remains partly disordered even in the bound state.<sup>56,57</sup>

Major changes are even more frequent in the other functional classes. They affect 7 of the 21 enzyme complexes with substrates or regulatory subunits. In thioredoxin reductase (1F6M), substrate binding affects the relative orientation of two domains in the enzyme; in HPr kinase (1KKL), it affects the orientation of a C-terminal  $\alpha$ -helix. Caspase-9/BIR3 (1NW9) is a remarkable system in which the interaction changes the quaternary structure. The active protease is a homodimer; it becomes inactive and monomeric in the complex, and BIR3 binds at the homodimer interface.<sup>58</sup> In complexes involving G-proteins, the G-protein contains "switch" segments that move, change secondary structure, or undergo disorder-to-order transitions when GTP or GDP binds, and often also when the G-protein interacts with another protein.<sup>59</sup> In the complexes involving receptors, the entire assembly of the receptor extracellular domains may be affected. For example, large domain rotations take place in the urokinase receptor, and the EPO receptor dimer rearranges completely when it binds erythropoietin.<sup>60</sup>

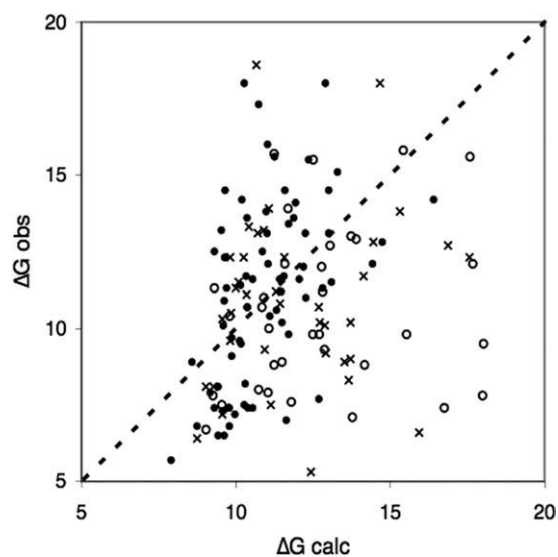
### Discussion

Since the early days of protein-protein interaction studies, relating structure to affinity has been a matter of concern to crystallographers as well as to biochemists and biophysicists.<sup>61,62</sup> However, these studies dealt mostly with individual systems, and the first attempt to associate binding affinities with a set of structures is due to Horton and Lewis,<sup>4</sup> who

collected from the literature 15  $\Delta G$  values, and showed that they could be fitted by summing contributions of the interface polar and nonpolar groups. The fit had just three adjustable coefficients, and it was remarkably good, yielding a linear regression coefficient  $R = 0.96$ , and a mean absolute difference of  $0.8 \text{ kcal mol}^{-1}$  between the calculated and observed  $\Delta G$  values. However, most of the data concerned protease/inhibitor complexes, and some of it was spurious. Trypsinogen/IleVal was given the same affinity for BPTI as trypsin, and a  $K_d$  was assigned to a hemoglobin S dimer (1HBS) that exists only in the crystal. Moreover, the authors noted that, whereas their formula fitted the  $\Delta G$  of trypsin/BPTI reasonably well, it was off by as much as  $10 \text{ kcal mol}^{-1}$  in the case of trypsinogen/BPTI. They attributed the discrepancy to the conformation change in trypsinogen, in line with Bode's analysis of the system.<sup>19</sup>

Later reports have used more diverse data sets, together with more elaborate models and more adjustable parameters.<sup>5–10</sup> None has achieved as good a fit to the data as Horton and Lewis,<sup>4</sup> and we can see at least two reasons for that. The first is the poor quality of the data sets, which contain many  $K_d$  values that are incorrect or associated with the wrong PDB entries, and others that cannot be traced to an actual measurement. The second reason is basic: the models rely on structural features of the complex alone. Thus, they represent the association reaction by its product, and ignore the reactants or the changes their structure may undergo. Whereas Horton and Lewis<sup>4</sup> had discussed the role of conformation changes, several recent reports do not mention them at all.<sup>8,10</sup> Audie and Scarlata<sup>5</sup> do, before stating that their contribution to  $\Delta G$  must be negligible. Their model fits very well a training set of 24 values, mostly for enzyme/inhibitor complexes, but it achieves about the same statistics as the other studies just cited ( $R = 0.73$ , root-mean-square  $\Delta G$  discrepancy =  $2.4 \text{ kcal mol}^{-1}$ ), on a more diverse control set of other 35 complexes.

A model of the association reaction based entirely on its final product is a plausible approximation if the components are known to behave as rigid bodies. The availability of the unbound structures in our benchmark allows us to state that this is incorrect except in a minority of cases, mostly antigen/antibody or enzyme/inhibitor complexes. In other systems, local, but significant, main chain movements take place at the interface, and one out four of the complexes displays large movements and/or disorder-to-order transitions. Their enthalpic and entropic costs contribute to the thermodynamic balance, lowering  $\Delta G$  by  $10 \text{ kcal mol}^{-1}$  in the case of trypsinogen binding BPTI. Figure 3 extends this remark to the whole dataset: when the main chain movements are of limited amplitude ( $I_{\text{rmsd}} < 1 \text{ \AA}$ ),



**Figure 3.** Conformation changes and binding free energy. Assuming  $\Delta G_{\text{calc}} = \alpha \Delta \text{ASA} + \beta$ , a linear regression of the observed  $\Delta G$ 's vs. the interface area  $\Delta \text{ASA}$  was performed on 70 complexes with  $I_{\text{rmsd}} < 1 \text{ \AA}$  (filled circles) excluding two (1BRS and 2PTC). The regression yields  $R = 0.54$  and a RMS discrepancy between  $\Delta G_{\text{calc}}$  and  $\Delta G_{\text{obs}}$  of  $2.4 \text{ kcal mol}^{-1}$ . When  $I_{\text{rmsd}} > 1 \text{ \AA}$ , the correlation with  $\Delta \text{ASA}$  vanishes, and 70% of the points are below the diagonal, meaning that observed  $\Delta G$  values are less than calculated ones (with  $P < 10^{-4}$ ). The average value of  $\Delta G_{\text{calc}} - \Delta G_{\text{obs}}$  is  $1.2 \text{ kcal mol}^{-1}$  for 39 complexes with  $I_{\text{rmsd}}$  in the range  $1\text{--}1.5 \text{ \AA}$  (crosses), and  $2.7 \text{ kcal mol}^{-1}$  for 35 complexes with  $I_{\text{rmsd}} > 1.5 \text{ \AA}$  (empty circles).

a  $\Delta G$  prediction scheme that only uses the interface area achieves performances similar to more elaborate empirical models used in the past; when the movements are large ( $I_{\text{rmsd}} < 1.5 \text{ \AA}$ ), the same scheme systematically overestimates the binding free energy by a large amount.

Prediction methods rely on experimental data to train and test procedures, and their performance critically depends on the quality and accuracy of those data. Collecting  $K_d$  from publications has proved to be a demanding task, validated in successive steps involving each of our groups. We did our best to ensure that the values we report are relevant to the PDB entries associated with them, and we expect them to be accurate to within a factor of 2–10 for  $K_d$ , or  $0.4\text{--}1.4 \text{ kcal mol}^{-1}$  for  $\Delta G$ . Nevertheless, they are valid only within a range of temperature, ionic strength and pH, and that range can be narrow, especially for pH. As a consequence, it makes little sense to model or predict a  $K_d$  to within better than an order of magnitude, unless one is also prepared to model its dependence on pH, and possibly also on ionic strength and temperature. This is particularly true of the low-affinity complexes, which were very few in early data sets, and are still under-represented, due in part to the difficulty in preparing crystals for structural studies.

In spite of some obvious limitations, like the absence of membrane proteins, the benchmark set of protein–protein complexes presented here covers a wide variety of functions and affinities. It should be an invaluable resource to computational structural biologists who attempt to predict binding affinity from structure, and stimulate the development of novel methods dealing with conformational changes. We found the collection of reliable binding affinity data to be a daunting task, and now hope to keep extending this dataset. As such, we call upon the community to provide binding affinity data, properly documented and when possible, associated with the deposition in the PDB of the structure of new protein–protein complexes.

## Methods

Information on the binding affinities was manually procured from literature. When it was ambiguous or incomplete, it was supplemented by personal communication with the authors. To maximize their reliability, the data were curated independently by each of the collaborating groups and compared to each other, so that all reported values have been checked in triplicate. Values from previously published sets,<sup>5,6,8,63–66</sup> including those collected for the complexes of the docking benchmark Version 3.0,<sup>11</sup> were used and controlled by checking primary citations. Data from publicly available databases (PINT<sup>40</sup> and ASEdb<sup>41</sup>) were also considered. Table S1 (Supporting Information) lists primary citations for all the final values.

Except when they were reported in the same publication as the structure itself, the affinity measurements have been done on protein samples, and possibly genetic constructs, that were different than the X-ray study, and under different experimental conditions. In cases where multiple sources were available, we selected those that most closely reproduced the content and the conditions of the crystal structure in terms of cofactors, ions, and pH, under which it was obtained. We then retained either the consensus value (the one that was found in more than one reference), or the value that we deemed to be most accurate, based on the measurement method or a personal communication with the authors. Most values were derived from direct physical measurements: SPR, ITC, and titration by fluorescence or other spectroscopic methods. They appear in publications in the form of equilibrium constants ( $K_d / K_a = 1/K_d$ ), or as the ratio  $K_d = k_d/k_a$  of rate constants issued from SPR and other kinetic measurements. For enzyme inhibitors,  $K_i$  values were assimilated to a  $K_d$  if they had been corrected for competition with the substrate and (when applicable) for slow binding kinetics.<sup>67</sup>

The temperature, pH, and experimental conditions were recorded when available. Taking the tem-

perature to be as stated (in 104 cases out of 144), or 25°C in other cases (room temperature, or no indication), we calculated the dissociation Gibbs free energy (in the  $c^\circ = 1M$  standard state) as:

$$\Delta G = -RT \ln (K_d/c^\circ)$$

For each complex in the benchmark, Supporting Information Table S1 reports the PDB entry codes and chain codes for the complex and its components, the  $K_d$  value and derived  $\Delta G$ , the reference to the publication reporting the measurement, the temperature, the pH, and method it used. The table also contains values of  $\Delta ASA$  and  $I_{\text{rmsd}}$ , consistent with the docking benchmark.<sup>13</sup>  $\Delta ASA$  is the accessible surface area (ASA) lost in the complex relative to its components in bound conformation; the ASA is calculated with program NACCESS<sup>68</sup> and a 1.4 Å probe.  $I_{\text{rmsd}}$  is the root-mean-square displacement of the C-alpha atoms of interface residues in the two partners, after the unbound and the bound interface residues have been superimposed by least-square.

A version of Supporting Information Table S1 with additional information on the affinity measurements and corroborating data is available online at <http://bmm.cancerresearchuk.org/~bmmadmin/Affinity>.

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