

# Detection of peptide-binding sites on protein surfaces: The first step toward the modeling and targeting of peptide-mediated interactions

Assaf Lavi, Chi Ho Ngan, Dana Movshovitz-Attias, Tanggis Bohnuud, Christine Yueh, Dmitri Beglov, Ora Schueler-Furman, and Dima Kozakov

#### ABSTRACT

Peptide-mediated interactions, in which a short linear motif binds to a globular domain, play major roles in cellular regulation. An accurate structural model of this type of interaction is an excellent starting point for the characterization of the binding specificity of a given peptide-binding domain. A number of different protocols have recently been proposed for the accurate modeling of peptide-protein complex structures, given the structure of the protein receptor and the binding site on its surface. When no information about the peptide binding site(s) is a priori available, there is a need for new approaches to locate peptide-binding sites on the protein surface. While several approaches have been proposed for the general identification of ligand binding sites, peptides show very specific binding characteristics, and therefore, there is a need for robust and accurate approaches that are optimized for the prediction of peptide-binding sites. Here, we present PeptiMap, a protocol for the accurate mapping of peptide binding sites on protein structures. Our method is based on experimental evidence that peptide-binding sites also bind small organic molecules of various shapes and polarity. Using an adaptation of ab initio ligand binding site prediction based on fragment mapping (FTmap), we optimize a protocol that specifically takes into account peptide binding site characteristics. In a high-quality curated set of peptide-protein complex structures PeptiMap identifies for most the accurate site of peptide binding among the top ranked predictions. We anticipate that this protocol will significantly increase the number of accurate structural models of peptide-mediated interactions.

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Key words: protein peptide interactions; FFT sampling; binding site detection; mapping; PeptiDB.

## INTRODUCTION

Much of the maintenance of a cell is accomplished by communication between proteins. Such communication may involve complex layers of regulation that are mediated by local features, such as changes in local concentrations of the partners or/and post-translational modifications. These interactions are often weak and transient, and tuned so that the threshold of biological downstream response may easily be manipulated. Peptide—mediated interactions, in which a short, linear stretch binds to its protein receptor, are well suited for such transient binding, and therefore extensively used. In higher eukaryotes, up to 50% of known interactions between proteins are indeed mediated by peptides. The support of the such communication of the such communication of the partners of the support of the such communication of the support of the support

The classical peptide–protein interaction involves (1) a short motif that is often embedded within an unstructured region, and (2) a peptide-binding domain with a defined globular structure. This interaction may occur between two distinct proteins, or within a protein, and

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the very competition between cis and trans interactions is often the very step that regulates protein function.<sup>4</sup>

One of the important sources of information about interactions is the structure of a protein-protein complex. This structure can be used as a starting point for the characterization and manipulation of an interaction. As an example, residues that are critical for an interaction may be identified using experimental or computational alanine scanning of interface residues.<sup>5–8</sup> Abolishment of an interaction by mutation of these critical residues may help identify the functional role of this interaction.<sup>9</sup> Finally, targeting of the interface of critical interactions by small molecules is gaining increasing importance in drug design, in addition to the traditional design of inhibitors of enzyme reactions. 10,11

Although the number of experimentally solved structures is increasing, the fraction of protein complexes among these remains very low, around 10-20%. 12 This calls for the development of approaches that identify a binding site on a protein structure, or even better, model the structure of a complex from the free monomers. Indeed, the field of docking, in which the structure of a complex is modeled from the structures of the free components, has significantly improved over the last 2 decades (see this CAPRI issue for some of the latest improvements).

Identification of the binding site on a protein structure is a first step toward the generation of an accurate structural model of an interaction. If crucial residues that mediate the binding of two partners can be identified, this has two important effects: First of all, experiments can be directed toward those residues and the functional effect of an interaction may be studied. Second, docking approaches may be focused on a specific interface patch.<sup>13</sup> For instance, we have previously developed a protocol that starting from a known binding site and an approximate peptide conformation within that site can accurately model the peptide-protein complex structure (FlexPepDock<sup>14,15</sup>), even without any detailed knowledge of the peptide structure within the binding site (ab initio FlexPepDock<sup>16</sup>). Thus, binding site identification allows to focus and to intensify the search to relevant sites, rather than wasting time in a global full docking search, which can also result in additional false positives.

Limited approaches have been proposed to identify peptide binding sites on proteins (e.g., Ref. 17-19). These use information both from the structures of the partners, as well as from the sequence.

PepSite identifies peptide binding sites on protein structures by searching for regions that match a spatial PSSM derived from known peptide binding protein receptor structures. 17 As such, it can not only identify the location of the peptide binding site but also suggests a sequence motif for the binding peptides. Consequently, information about the actual peptide-binding partners is also provided. Another recently published approach uses the BRIX database of interacting fragments to predict the structure of peptide-protein complexes starting from a peptide sequence and a solved receptor structure. 19

As for peptide binding sites, these existing methods perform well mainly on known binding sites, such as WW, SH3, and kinase domains, but less well on nonstandard peptide-mediated interactions. Thus, new tools are needed to address this problem.

Here, we suggest an approach based on the observation that protein functional sites, including peptide binding sites, also bind small organic molecules of various shapes and polarity, as observed by nucleic magnetic resonance (NMR)<sup>20</sup> and X-ray crystallography experiments.<sup>21</sup> FTmap<sup>22</sup> is a direct computational analog of the above experimental approaches. This protocol is based on the successful FFT-based docking protocol with statistical potential.<sup>23</sup> We have recently reported application of this approach to ligand binding site identification<sup>24</sup> and druggable protein-protein interaction sites,<sup>25</sup> demonstrating broad range commonality of principles of molecular recognition.

In this study, we have calibrated a protocol to detect peptide binding sites on protein structures. For this, we have adapted the mapping protocol to the identification of peptide binding sites. Two key differences distinguish peptide site prediction from ligand binding sites detection: (1) Prediction requires knowledge of the domain critical for peptide interaction: In contrast to proteins that bind ligands at one key site, multidomain proteins can have multiple peptide regulatory sites on each of the domains and therefore focusing on one particular domain is recommended; (2) Peptides do not bind to inner buried sites of the proteins, which could well be ligand binding sites, and therefore these internal sites need to be removed. We show that with this tailored protocol, we can identify accurately the binding site of a peptide within the top-3 sites in 19/21 (90%) of a benchmark set of curated peptide-protein complexes. We validate the approach on a set of peptide-protein structures that were released after calibration of the protocol: for (78%) of the structures we reliably identify the peptide-binding site. In addition to the robustness of our protocol, our results also highlight features that characterize specifically peptide-protein complex structures, and finally allow us to identify yet outstanding challenges in the modeling, and also the experimental identification of peptide-mediated interactions.

## **METHODS**

## Compilation of training and test sets for protocol calibration

In a previous study we have studied the special character of peptide-protein complex structures. For this aim, we collected PeptiDB, a set of 103 peptide-protein complexes that represent a set of different known peptide-mediated interactions (less than 70% sequence identity among different protein receptors), and a more restricted set of 61 complexes in which no two proteins share the same fold (according to CATH classification) (See Ref. 26 and Supporting Information Table S1 therein).

In this study, our aim is to identify the peptide binding site on the protein surface. In order to prevent bias to bound sites on a protein structure, we needed to verify that the surface accessible to the peptide for binding on the receptor is indeed not occupied by any other protein or ligand. We therefore went over the original set and checked each protein-peptide interaction individually for the following features:

- 1. Availability of the free receptor structure. We filtered for cases in which the corresponding free conformation has been solved by X-ray crystallography at reasonable resolution (<2.7Å), for a protein with full (>98%) sequence identity.
- 2. No bound ligands or proteins on the surface of the receptor. We verified that no ligands are bound to the protein surface, in particular not to the binding sites (e.g., for the PeptiDB entry 1T4D (mdm-x), we could not find a corresponding structure of the free mdm2 molecule without any ligand that had been solved by crystallography, and therefore we removed it from our set).
- 3. No crystal contacts that stabilize the binding site. Inspection of the symmetry mates in the solved crystal structures revealed that in certain cases the binding site of the free conformation binds to a peptide stretch of a copy of the protein in the crystal lattice (see for example PeptiDB entry 1DDW, the evh1 domain in the homer protein). In these cases, the peptidebinding site might have been arranged in a similar way as the bound conformation, and this might bias our protocol.
- 4. Biological unit of protein. In order to simulate accurately the accessible surface to the peptide, we included in the dataset the biological unit of the protein, for example, a homodimer (see for example PeptiDB entry 2DS5 of the CLPX protease Zn binding domain).

The resulting set for assessment is detailed in Table 1A. To assess the robustness of this protocol, we compiled an additional set by extracting peptide-protein complex structures that were released to the Protein Data Bank (PDB<sup>12</sup>) after the development of PeptiMap (between the dates 1.1.2013 and 8.4.2013) and filtered according to the same guidelines as detailed earlier. Only protein receptor structures not structurally similar to the entries in the training set were retained (i.e., distinct CATH domain<sup>27</sup>). This validation set is detailed in Table 1B.

#### Definition of the functional unit that will be mapped

Similar to its experimental counterpart, Structure-Activity-Relationship (SAR)-NMR, <sup>28</sup> our computational solvent mapping approach is aimed at targeting distinct "physiological units," namely globular domains that form a stable unit and together act as a receptor. As an example, a tight dimer receptor will be defined as one physiological unit to map. In contrast, when a protein is composed of distinct domains that can move one relative to the other, we expect each to act as an individual binding site to a peptide, and therefore each should be treated as a distinct physiological unit. On the basis of this rationale, we have devised the following rules by which we define the individual units that are separately mapped:

- a. For "tight" multimers (ratio of buried surface area of the monomer >0.2, see for example 1GY7), the individual unit is defined as the full multimer.
- b. If a protein consists of more than one domain, all of the same class (i.e., repeated domains), we do not split the protein into individual domains, but rather treat it as one individual unit.
- c. If the protein consists of different domains, split these into individual domains and perform solvent mapping separately on the domain that is known to be critical for the interaction (if such information is not available, each domain should be screened separately).

#### Separation of receptor structure into distinct domains

In order to map the fragments onto individual domains, the protein receptor structure was decomposed into these domains based on CATH domain classification (v3.4). In cases where no CATH classification was available, we identified the most similar CATH domain using sequence alignments.

## Detailed outline of protocol

The protocol consists of a series of steps described below. First, the "physiological unit" of the protein receptor is defined (as detailed earlier). The next steps (2-7) are identical to the FTsite protocol used to identify protein and ligand binding sites.<sup>24</sup> The final steps 8–10 include the removal of sites involved in domain interactions, the merging of adjacent clusters, and finally the filter of internal sites not accessible to the peptide. Here, we summarize the protocol shortly, and where appropriate highlight the changes specific to peptide binding site identification that were incorporated into PeptiMap.

Step 1: Selection of "physiological unit". Decomposition is based on annotation of protein domains according to CATH (the rules for decomposition are outlined above and more detailed in the Results section).

**Table 1** Benchmark and Validation Sets Used for Optimization and Evaluation of PeptiMap Peptide Binding Site Identification

			A. Initial calibra	A. Initial calibration benchmark (Extracted from PeptiDB <sup>26</sup> , <i>n</i>	om PeptiDB $^{26}$ ; $n = 21$ )			
Name	punoq	punoqun	Peptide sequence	Cath domain	domains/multimer	treatment	rank of hit	Pepsite2 <sup>18</sup>
dystrophin (WW)	1EG4A	1EG3A_1	NMTPYRSPPYVP <sup>a</sup>	2.20.70.10 ( <sup>b</sup> ;	1 of 4 different domains	split: only first domain	2,3	6,8,10
sh2a1 ( <b>SH2</b> )	1D4TA	1D1ZA	KSLTIYAQVQK	3.30.505.10			- c	 -
erbb2 ( <b>PDZ</b> )	1MFGA	2H3LA	EYLGLDVPV	2.30.42.10			1 -	<u> </u>
wdr5 ( <b>WD40</b> )	2H9MA	2H14A	A <u>r</u> tkot	2.120.10.80	central hole not acces- sible to peptide	mask internal sites	က	1,3,5,6
usp7 cyclophilin	2F0JA 1AWR	2F1WA 2ALFA	GAR <b>AHSS</b> HAG <u>P</u> IA	NA <sup>d</sup> (β-sandwich) 2.40.100.10			1,2	1–6 1–10
p97 IN-glycanase traf2	ZHPLA 1CZY	2HFJA 1CA4A	ace <u>-P</u> û0 <u>A</u> TDD	NA (p97) 2.60.210.10	weak homotrimer (ABC)	use monomer <sup>c</sup>	– ო	1 1
i-ap1 gga1	1JD5A 1JWG_AC	1JD4A 1JWFA	AIAYFIPD <u>Dedl</u> lhi	1.10.1170.10 1.25.40.90	weak homodimer (AB) weak homodimer (AC -	use monomer use monomer	1,2 2	4
ntf2	1GYB_AB	16Y7_AB	FSF	3.10.450.50	in bound structure) tight homodimer	unit: 2 chains		1
Cıpx calpain small subunit	20.38_AB 1NX1A	2025_AB 1ALV_AB	ALKVVK DAIDALSSDFT	NA* (cipx) 1.10.238.10	tignt nomodimer, z sites tight homodimer central hole not accessible	unit: 2 chains unit: 2 chains <sup>c</sup>	<b>7</b> ' –	1 1
					to peptide			
ap2	2VJ0A	1B9KA_1 1B9KA_2	PKGWVTFE Fednevp	2.60.40.1030 3.30.310.30	2 domains	split: both have peptide bound	1 m	1 1
pim1 kinase	2C3I	2J2IB_2	<u>K<b>r</b></u> rrhpsg	(3.30.200.20) 1.10.510.10	2 domains	split: only second used	2	ı
transterase domain								
cdk2 cyclin	2CCHB	1H1RB	HTLKGRRLVFDN	1.10.472.10 x2	2 same domains	do not split	က	1,6
trypsin	2AGEX 1RX7	1UTNA 1RW7A	Sin-AAP <b>R</b> KSTOATI <sup>E</sup> B\WE	2.40.10.10 x2 3 70 10 10	2 same domains	do not split	- 2	9
Endothiapepsin	1ER8	4APEA	PFHLLV	2.40.70.10	2 same domains	do not split	1,2,3	1-6
			B. Validation set (	B. Validation set (compiled from recent PDB releases 1–4/2013; $\it n$	eleases 1–4/2013; $n=9$ )			
Name	punoq	punoqun	Peptide sequence	Cath domain	domains/multimer	treatment	rank of hit	Pepsite2
γ2 adaptin Ear domain	2YMT	4BCXA	EWGP <u>WV</u>	2.60.40.1230	1 domain		က	l dilks
в сор WD40	4J73	3MKQA	EAKKI <u>v</u>	2x 2.130.10.10 + additional domain (based	3 domains central hole not accessible to	split, only first domain (2–300); site 1	2,3	2,3,5,6
RADa	4B3B	4A74A	ace <u>F</u> HT <u>A</u>	on 1VYH) 3.40.50.300 (based on	peptides weak homodimer	masked use monomer	2,3	ı
FKBP35	4ITZA	3NI6A	sinALPFnit	1PZN) 3.10.50.40 (based on	weak homodimer	use monomer	_	1-10
jnk1 kinase	зупн	3ELJ	PKRPTTLNLF	1KT0A) (3.30.200.20); 1.10.510.10	2 domains	split: only second	1	1–10
transferase domain						domain		

Name	punoq	punoqun	Peptide sequence	Cath domain	domains/multimer	treatment	rank of hit	Pepsite2 ranks
HIF alpha	4B7E	2W0X	EVVKLLLEHGADVLAQD	2-305; 1.10.287.1010		split: only first domain	1,3	1,4,7-10
MLH1 mut alpha c-term	4FMNA	4E4WA	VRSKYFK	NA <sup>d</sup>	multidomain	too big without splitting <sup>c</sup>	–No split information available	I
demethylase	4FWF	4FWE	ART <u>M</u> OTARKSTGG	(domain A02: 280–405 and 522–836, based on 2V1DA)	multidomain	split: only second domain	_	I
ck2 kinase	41B5	3800	GCRLYGFKIHGCG	3.30.200.20; (1.10.510.10)	2 domains	split: only first domain	33	ı

Peptide residues accurately mapped by fragments are highlighted in bold and underlined; Peptide residues with area covered by fragments are highlighted in bold but not underlined XX: unassigned domains.

Better prediction obtained with SCOP classification (1CA4: 1,2 instead of 3), or domain parser (1ALV)

Step 2: Grid-based sampling of the protein surface with FFT. All bound ligand water molecules and other ligands are removed prior to the calculations. We then sample the protein surface for 16 small molecule probe types.<sup>22</sup> This is done using exhaustive sampling with grid-based Fast Fourier Transform (FFT) (109 docked probe positions). The best 2,000 poses with the lowest energies for each probe type are retained.

Step 3: Post-FFT clustering to discard spurious probe clusters. For each probe type, the 2000 retained poses are clustered using a simple greedy algorithm. We select the lowest energy pose as the center of the first cluster, and add all poses within 4Å center-to-center distance from it as cluster members.<sup>29</sup> All clustered poses are removed, and we repeat the same steps to form the second and then the subsequent clusters until all poses are clustered. Clusters with less than 10 probes are removed, and the six largest clusters are retained for further analysis.

Step 4: Minimization and rescoring. The energy of each retained protein-probe complex is minimized using the CHARMM<sup>30</sup> potential with the Analytic Continuum Electrostatic (ACE) model representing the electrostatics and solvation terms as implemented in version 27 of CHARMM. The algorithm uses the polar-hydrogen-only parameter set from version 19 of CHARMM. The energy minimization is performed using a limited memory Broyden–Fletcher–Goldfarb–Shanno (L-BFGS)<sup>31</sup> method in which heavy atoms of the protein are held fixed, whereas the polar hydrogen atoms of the protein and all atoms of the probes are free to move. Poses with positive energies after minimization are discarded.

Step 5: Generating consensus clusters. Following the energy minimization we recluster the resulting probe poses. As in Step 2, we select the lowest energy pose as the center of the first cluster, but use 4 Å full-atom pairwise RMSD as the clustering radius. After all probes are clustered and clusters with less than 10 members are discarded, the clusters are ranked on the basis of the Boltzmann averaged energy, and the six lowest energy clusters are retained for every probe type. Consensus clusters are generated by grouping probe clusters with cluster centers within 4 Å. The centers of the resulting consensus clusters are fixed, and the probe clusters are redistributed such that each cluster center is closer to the center of its own consensus cluster than to the center of any other consensus cluster. Consensus clusters that overlap with an integral element of the intact protein such as heme are discarded. A consensus cluster is considered to overlap with a cofactor if their volume overlap exceeds 80% of the consensus cluster.

Step 6: Ranking consensus clusters. The algorithm ranks the consensus clusters by the number of nonbonded contacts between the protein and all probes of the consensus cluster. A residue of the protein and a probe are considered to be in contact if any atom of the residue is less than 4 Å from any atom of the probe. A residue is considered to be in contact with a

consensus cluster if it is in contact with any of its probes. After selecting the contact residues for a consensus cluster we re-evaluate the number of contacts by adding also interactions with probes that are within 4 Å but are not part of the original consensus cluster. The resulting numbers are normalized using the overall number of contacts for all probes, and used for ranking the consensus clusters.

Step 7: Identification of putative peptide binding sites. To identify the putative binding site, the algorithm first selects the consensus cluster with the highest number of contacts. This cluster is then expanded by adding any neighboring consensus clusters if the center of any of its probe is closer than 3.5 Å to the center of any probe in the consensus cluster. The protein residues that are within 4 Å of the expanded consensus cluster constitute the top prediction of the binding site. The first consensus cluster is then removed, and the procedure is repeated using the next consensus cluster with the highest number of contacts to identify lower ranked predictions of the peptide-binding site.

Step 8: Discard sites located in nonaccessible regions in the protein. Sites at the domain interface (in cases of proteins that were split into individual domains): If a putative site clashes significantly with secondary structures ( $\alpha$ -helices or  $\beta$ -sheets) on the partner domain(s), the site is discarded. A putative site is discarded if the site is within 3.0 Å of one or more secondary structureassociated (helices or sheets) amino acids on partner domain(s) as defined by PyMol. This removes sites that are involved in domain-domain interactions.

Step 9: Expand final sites that are retained. Each of the sites is expanded by adding to the site probe cluster representatives that are not already part of the site but are closer than 4.0 Å (atom-atom distance) to probe cluster representatives that are already part of the site. Probe cluster representatives that are already members of other sites are not used and, therefore, the sites are only expanded and not cojoined.

Step 10: Remove inaccessible sites within the protein core: Whereas the small molecules that are used to probe the protein surface may access internal cavities due to their small size, peptides are larger and therefore cannot reach these voids. At the same time such sites are attractive to probes as they provide much larger contact areas to a small molecule than the surface (and indeed they are good ligand binding sites). To adequately analyze peptide binding surface, such sites should therefore be excluded from mapping. In order to identify internal ligand binding sites, we build 100 rays uniformly covering a sphere<sup>32</sup> from the center of the site, and identify which of those contact the protein (a contact is defined if a ray passes within 2 Å from the center of any atom of the receptor). A binding pocket is considered internal if 80% of the rays contact the protein. Internal sites are masked and steps 2-10 are repeated.

## Assessment of performance: criteria for binding site identification

As peptide binding site identification criteria have not yet been established, we have employed accepted criteria used for ligand binding site identification.<sup>33</sup> This criterion requires the geometric center of the predicted ligand-binding site to lie within 4.0 Å of any peptide atom. This allows easy comparison to other approaches. Residues mapped by PeptiMap are shown in bold in Table I, and residues for which side chain atoms have been accurately mapped at the same spatial position (i.e., within 2 Å are underlined).

## **RESULTS**

## Compilation of curated benchmark set of peptide-protein complexes for peptide binding site prediction

In order to allow the objective assessment of peptide binding site prediction, we compiled a set of protein structures of a free receptor that do not contain any other molecule at the peptide binding site (i.e., no other bound ligands, and no crystal contacts, see Methods section for more detail). We then parsed the structures into domains and identified their biological units. On the basis of these, we defined the receptor structure and the surface to mapped for peptide binding sites. The same procedure was repeated on a second set compiled after this calibration (see Methods). The total of 30 peptide–protein complexes are detailed in Table I (benchmark of 21 interactions and additional validation set of nine interactions).

We would like to note that while the optimization set, and more so the test set, are rather small, they are nonredundant and therefore do represent a wide variety of different peptide binding domains. More importantly, considerable efforts were made to verify that these sets are clean and not biased to any bound conformation. In a real-world scenario the available template structure might well bind a symmetry mate or a ligand in the binding site-thus PeptiMap is expected to perform even better.

## Adaptation of FTsite to prediction of peptide binding sites

Computational fragment mapping can accurately identify small ligand binding sites,<sup>24</sup> as well as locate druggable protein-protein interfaces. 25 Peptides lie in between these two cases. In contrast to ligand binding sites, peptide binding sites tend to be more shallow and the pockets to be smaller. In contrast to protein binding sites, peptides tend to bind to smaller regions and with usually weaker affinity. We wanted to optimize PeptiMap to specifically identify peptide-binding sites (noting that such modifications might improve predictions of binding sites of larger ligands as well). Two major modifications were necessary to provide a robust protocol for peptide

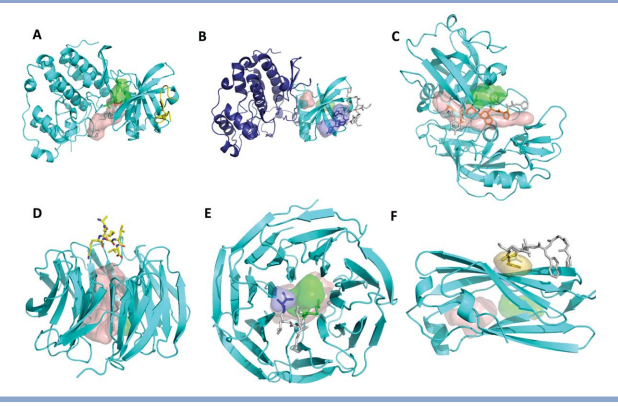


Figure 1

Examples of predictions of peptide binding sites using PeptiMap. The same color-coding scheme is used in all figures: The receptor is shown as a cartoon colored in cyan, while the different predicted sites are shown in surface representation (top-ranking site-wheat; site 2-green; site 3-dark blue). The peptide is shown in stick representation and colored according to the predicted binding sites (residues that do not contact any predicted site are in light grey). Pdb IDs of the apo and bound structures are indicated in parentheses. Predictions were made on the apo structures; bound structures are for validation purposes only. (A,B) Mapping on single domains improves peptide binding site prediction on ck2 kinase (apo: pdb ID 3BQC chain A): (A) mapping on the full structure identifies merely a known ligand binding site (pdb ID 3U4U), while (B) mapping on the n-terminal kinase domain only identifies the peptide binding site (pdb ID 4IB5; the transferase domain not mapped here is shown in dark blue). This highlights how implementation of segmentation into single domains may improve in particular peptide-binding site predictions (while small ligands can be identified also on the full structure). (C) Example of accurate PeptiMap prediction of the peptide binding site covering a substantial part of the peptide: Endothiapepsin peptide binding site prediction identifies the peptide binding site of all but one residue (apo: pdb ID 4APE chain A; bound: IER8). (D,E) Masking of internal ligand binding sites improves peptide binding site prediction on coatomer b subunit. (D) In the original prediction, mapping of the receptor structure (pdb ID 3MKQ chain A) identifies mainly peptide-inaccessible sites within the hole of the WD40 domain, but not the peptide binding site (pdb ID 4J73). (E) When entrance into the inner cavity is blocked, PeptiMap identifies correctly the location of the peptide-binding site. (F) Example of target failed by Peptimap. In case of the AP-2 complex subunit alpha (CATH domain 2.60.40.1030; pdb ID 1B9K), the peptide binding site (from pdb ID 2VJ0) is not identified by the top 3 predictions, only by prediction ranked 4th (in yellow).

binding site location on protein structures, namely (1) the decomposition of protein structures into physiological, functional units, and (2) filtering out internal ligand binding sites not accessible to peptides.

#### Decomposition of protein structures into physiological, functional units

Assuming that a peptide will bind to an organized binding site, we decompose the protein structure into independent parts that represent the stable functional unit a peptide might encounter. Splitting a protein structure into individual domains prevents the identification of short-lived crevices at the boundary of the domain interface as peptide binding sites, and reduces the surface to be sampled [see Table I, and Fig. 1(A and B) for the example of ck2 kinase, pdb ID 3BQC].

In contrast, tight homo-multimers (e.g., Clpx, pdb ID 2DS5), as well as repeated same domains within a protein (that in general also form tight interactions) are merged into one functional unit for mapping [e.g., Figure 1(C) that shows Endothiapepsin chain A that is composed of two identical domains according to the CATH classification].

# Filter out internal binding sites

Fragment mapping will identify among others also ligand-binding sites within proteins that are not accessible to the larger peptide ligands. Removal of such sites using ray tracing (see Methods section) improved predictions of peptide binding sites of WD40 domains that contain a hole at the center of the protein not accessible to peptides. Figs. 1D and E show how ranking is significantly improved by masking inaccessible internal ligand binding sites in the protein cop b (pdb ID 3MKQ chain A), and similar results were also observed on additional cases of known internal ligand binding sites (results not shown).

#### PeptiMap accurately locates the peptide-binding site on most of the receptor structures in both the benchmark and validation sets, and compares very favorably to other approaches

Using this streamlined protocol, PeptiMap identified 10 out of 21 (50%) peptide binding sites in the benchmark as the top-ranking prediction. More importantly, it failed only in two cases to identify the site among the top-3 ranking predictions (19/21 success rate; 90%). In the independent subsequent validation, a lower performance was observed for top-ranking sites: 3/9 (33%) binding sites were top-ranked. However, again only two cases were not identified among the top-3 ranked predictions (7/9 successes; 78%). Details of performance for each of the proteins are given in Table I. These results demonstrate the general applicability of the protocol to a range of different peptide-protein complexes, involving many different folds, as well as many different functional classes.

The predicted binding sites are in many cases very accurately mapped. Figure 1(C) shows a particularly successful prediction: for Endothiapepsin, the accurate location of six out of seven peptide residues is identified by PeptiMap. Overall, 1-6 peptide residue positions (median of 2) are identified by PeptiMap (these residues are highlighted for each peptide in Table I). It should be noted that this covers a significant part of the peptide residues that directly contact the receptor.

We compared our protocol with other available approaches. PepSite is the only approach that can be tested and validated via a server. The coverage of this approach is rather restricted (due to limited availability of enough structures to create the structural PSSMs). We ran PepSite (version2<sup>18</sup>) on the same dataset in order to compare the two approaches. PepSite identified only 6 out of 21 peptide binding sites in the benchmark set within the top1 predicted sites (same results for top-3 assessment). The corresponding performance for the validation set of nine cases was three top-ranking predictions and four predictions ranked 1-3.

The recently published approach based on the BRIX database of interacting fragments 19 reports results for a set of protein-peptide complexes. Among these we could find predictions for two of the proteins assessed here: for p97 N-glycanase (2HPJ) this approach failed to identify the binding site, whereas an acceptable prediction was reported for PCNA (1RWZ), albeit not top-ranked.

# **DISCUSSION AND** CONCLUSIONS

PeptiMap is a new, accurate and robust approach for peptide binding site detection on protein receptor surfaces. It is based on the successful computational fragment mapping approach previously applied to ligand site prediction<sup>24</sup> and the detection of druggable protein interactions.<sup>25</sup> Overall the results presented here are promising and indicate that automated and efficient prediction of peptide binding sites on proteins is coming of age. Not only is the peptide binding site on the receptor surface identified for most of the cases among the top-3 ranking structures (26/30) but also the predictions are accurate and identify many of the important peptide binding residues. Furthermore, our results here suggest that Pepti-Map clearly outperforms other available approaches, thanks to its general applicability and accurate prediction.

In the following we shortly assess the challenges that need to be addressed to further extend performance of PeptiMap and suggest several ways to do so, based on the constraints of this approach and the cases where it still fails [see e.g., Fig. 1(F) for a peptide-binding site identified only by the prediction ranked 4<sup>th</sup>].

#### Possible strategies to improve Peptimap

- 1. Focused mapping We have shown in several examples that once a site has been identified, focused local mapping can improve the coverage of the site, and provide general guidelines regarding peptide sequence preference.<sup>7,34</sup>
- 2. Improved definition of functional units: Preliminary results indicate that performance can significantly be affected by how the functional units are defined, in particular how the individual domain boundaries are determined. While CATH-based domain definition provides overall adequate functional units, in some cases we noticed that different definitions can dramatically improve performance (see Table I). In particular for large proteins, when CATH provides no domain definition or does not split the protein into distinct domains, PeptiMap may fail to identify the peptidebinding site (e.g., pdb ID 4E4W chain A and 1ALV chain B, see Table I). In such cases, domain mapping based on alternative tools such as domainparser,<sup>34</sup> or on visual inspection could define a subdomain that is useful for peptide binding site detection (results not shown).

## Combination of PeptiMap with other approaches to characterize structure and specificity of peptide-protein interactions

Currently, PeptiMap predicts the location of peptide binding sites, but does not provide any information about actual structure of the peptide within this site nor about possible sequences for a binding peptide. In order to proceed to a full model of the peptide-receptor complex, PeptiMap predictions could serve as input for

peptide-protein docking protocols. 13,36 We plan to incorporate PeptiMap into a scheme for FlexPep-Dock 14,16 that will allow for full ab initio prediction of peptide-protein complex structures starting from a given peptide sequence and a free receptor structure.

In addition, whereas PeptiSite and the BRIX-based approach are less general, they do provide more information in their prediction, such as sequence-specific binding site identification, as well as an approximate structure of the peptide-protein complex. Therefore, combining information from different protocols that use rather complimentary approaches will ultimately improve our knowledge and understanding of peptide-mediated interaction and their structural basis.

#### Implications for globular protein-protein docking

In a previous study we have suggested that a significant fraction of globular protein-protein interactions is mediated by one linear, peptidic stretch that contributes most of the binding energy (e.g., a dominant loop at the interface).<sup>37</sup> Moreover, this study indicated that when bound to the protein partner, this peptide tends to adopt a structure that is very similar to the one adapted within the protein context. Consequently, PeptiMap could be used in a more general way to identify on a protein surface where such dominant peptidic stretches would bind. Although FTmap was shown to identify druggable sites on protein interfaces, 25 PeptiMap is expected to be very useful for the design of specific peptide-derived inhibitors of protein interactions.

This study demonstrates the strength of nonbiased, ab initio prediction protocols for finding molecular recognition sites of peptides. The general applicability of such an approach will substantially contribute to improved characterization of a range of peptide-mediated interaction, and provides thus a good starting point for structure-based characterization of biological interaction and function. A server is under development to make PeptiMap generally available (and the PeptiMap software is freely available to academic users upon request).

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