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Docking approaches for modeling multi-molecular assemblies
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Computational docking approaches aim to overcome the limited availability of experimental structural data on protein–protein interactions, which are key in biology. The field is rapidly moving from the traditional docking methodologies for modeling of binary complexes to more integrative approaches using template-based, data-driven modeling of multi-molecular assemblies. We will review here the predictive capabilities of current docking methods in blind conditions, based on the results from the most recent community-wide blind experiments. Integration of template-based and \textit{ab initio} docking approaches is emerging as the optimal strategy for modeling protein complexes and multimolecular assemblies. We will also review the new methodological advances on \textit{ab initio} docking and integrative modeling.

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Introduction
Protein–protein interactions are key for the majority of biological functions. Proteins can form highly specific transient or permanent complexes that range from binary pairs to multi-molecular assemblies, often involving other biomolecules. A detailed structural knowledge of such complexes at atomic level would improve our understanding of biological processes and facilitate intervention for biomedical and biotechnological purposes. For example, recently reported structural data on the dynamic assembly formed by the SARS-CoV-2 trimeric spike protein and the cell receptor ACE2 are key to understand the molecular mechanisms of the virus infectivity and can be essential for the development of new vaccines and therapeutic candidates against COVID-19 \cite{1,2,3}. However, structural data is available for only a small fraction of the protein interactome. For instance, the total number of protein–protein interactions in human is estimated to range from 130,000 \cite{4} to 650,000 \cite{5}, but less than 7000 of these interactions have available 3D structure (Interactome3D, 2019.1 version) \cite{6}. In this context, computational docking approaches aim to overcome the limited availability of experimental structural data. Since the first reported protein-protein docking algorithms in the early 90’s, based on Fast Fourier Transform (FFT) sampling \cite{7}, the methodological developments have mostly focused on \textit{ab initio} docking of binary complexes, starting from the structure of the unbound components. Some of the most popular methods are FTDock \cite{8}, ZDOCK \cite{9} or MolFit \cite{10}. The method HEX \cite{11} and later FRODOCK \cite{12} used polar Fourier correlations to accelerate docking calculations. Other different approaches using stochastic search based on global-energy optimization are ICM-DISCO \cite{13,14}, RosettaDock \cite{15}, HADDOCK \cite{16}, or SwarmDock \cite{17}.

With the increasing availability of complex structures, in recent years attention is focused on template-based structural modeling of complexes, based the standard principles of homology-based modeling. The term template-based docking (as opposed to \textit{ab initio} docking) is specifically used when a model is built by superimposing the structures (or models) of the unbound subunits onto the corresponding subunits of a template complex structure \cite{18}. One advantage is that template-based modeling can be applied to multi-molecular complexes, not just to binary complexes as \textit{ab initio} docking. In addition, it has been suggested that templates are available for the large majority of cases in which interacting subunits have structural information \cite{19}. However, the general availability of good-quality templates that could be reliable used for template-based predictions seems much lower \cite{20}. Actually, for the majority of known interactions, only templates with remote homology are available \cite{4}, for which direct application of template-based methods leads to poor predictions \cite{21}. Modeling multi-molecular assemblies implies additional challenges. For instance, some of the interfaces might not have available templates, in which case, we could model them by \textit{ab initio} docking, in combination with restraints from evolutionary data or from available experimental information. Another challenge is to identify the relevant oligomerization state of the assembly when is different from that in the template.
[22], in which case, alternative orientations provided by 
_ab initio_ docking can be very helpful. Modeling the 
conformational variability of the assembly components 
imposes an additional difficulty. Indeed, directly 
taking the structure of a given subunit in another context 
(e.g. unbound state, different assembly or alternative 
oligomerization state) might lead to inaccurate models. 
For this, it can be useful the application of protein-protein 
docking and associated procedures, such as energy 
scored, minimization, or flexible refinement.

We will review here the predictive capabilities of current 
protein-protein docking methods in blind conditions, 
based on the results from the most recent CASP [23**] 
and CAPRI [24*] experiments. These tests show that 
combination of template-based and _ab initio_ docking 
approaches is emerging as the optimal strategy for modeling 
protein complexes and multimolecular assemblies. We 
will also review the most recent methodological 
ovancements on _ab initio_ docking, and new approaches for 
the inclusion of experimental information and integrative modeling.

**Predictive capabilities of computational docking: the state-of-the-art**

_Ab initio_ computational docking can provide acceptable 
models within the top 10 predictions in up to 40% of the 
cases, according to reported evaluation studies of different 
methodologies in current protein–protein docking 
benchmark version 5.0 [20*,25,26].

Traditionally CASP has been focused on the prediction of the 
structure of individual proteins. However, very often 
proteins are found as oligomeric assemblies, which adds 
complexity to the modeling effort. To evaluate the 
applicability of docking methodologies for the prediction of 
protein oligomeric assemblies, the last three CASP 
editions included a CASP-CAPRI joint experiment 
focused on multimeric assemblies, which are independently evaluated by CASP and CAPRI communities. 
The recent CASP13-CAPRI challenge comprised a total of 
20 protein oligomeric assemblies, including 14 homo-
complexes and 6 hetero-complexes, which could be classified into 15 dimers and 5 multimeric assemblies 
[23**]. In the 9 ‘easy’ targets, there were good structural 
templates for the (partial or full) assembly, while for some of the remaining 11 ‘difficult’ targets, it was possible to 
find remote templates for part of the assembly. The 
availability of templates in each case is critical to explain 
the predictive success of the groups. Focusing on the 
results for the top 10 predictions (to facilitate comparison with the reported performances of different docking methods in the literature), the best-performing group submitted acceptable (or better) models for 13 targets (65% of the cases) (Figure 1). In the ‘easy’ targets, the 
best-performing group submitted acceptable models for all these cases, while in the ‘difficult’ targets, the 
best-performing group submitted acceptable models for only 4 of such targets (36% of the cases). Regarding 
the quality of the models, high-quality models [23**] were submitted by any group in 78% of the ‘easy’ targets 
(with template), but only in 9% of the ‘difficult’ targets (no template).

On the other side, the recent 7th CAPRI edition showed 
more heterogeneity in its targets, comprising 8 protein-
protein, 3 protein-peptide, and 5 protein-oligosaccharide 
complexes, all hetero-oligomers (except for a homodecamer), which could be classified in 10 dimers and 
6 multimeric assemblies [24*]. The actual number of 
evaluated targets was 19, because some of the interfaces 
in these multimeric assemblies were considered as independent targets. There were structural templates 
for a total of 13 target interfaces (6 protein–protein, 2 protein–peptide, and 5 protein–saccharide). This was 
determinant for the overall predictive success of the 
groups as well as for the quality of the predicted models. 
Overall, the maximum number of target interfaces successfully predicted by a single group was 13 (i.e. 
success in 68% of the cases) (Figure 1). But in cases with 
no available template, the best-performing groups submitted acceptable models for only 2 target interfaces (i.e. 
success in 33% of the cases). Regarding the quality of the 
models, high-quality models [24*] were submitted by any group in 31% of the ‘easy’ targets (with template) and in 
17% of the ‘difficult’ targets (no template). The 7th CAPRI edition showed that _ab initio_ docking in cases 
for which there is no available template is still highly 
challenging, and progress is actually coming from the efficient procedures to combine template-based 
modeling and other docking methodologies.

**Combination of template-based and _ab initio_ docking**

The CASP and CAPRI experiments show that template-
based modeling approaches are clearly the tools of choice 
when one can use templates of sufficient quality. 
However, very often only remote templates are available, 
which might not be good enough to provide reliable 
models, as above discussed [21]. In unclear situations, 
a relevant question is which method to choose, or how to 
efficiently combine these protein-protein docking 
approaches depending on each specific case [20*]. This 
is even more relevant when modeling multimeric 
complexes, in which some interfaces might be modelled 
based on homologous structures, while others would need _ab initio_ docking, as above mentioned. An updated 
version of the InterEvDock2 server [27**] can perform 
template-based docking or _ab initio_ docking with 
evolutionary constraints, depending on the case. But 
the question is still open about how to efficiently combine 
template-based and _ab initio_ docking when reliability of 
the template is unclear. We can obtain some hints from 
the recent CASP and CAPRI experiments.
In the recent CASP13-CAPRI joint assembly prediction experiment, one of the most efficient approaches was that of Fernández-Recio, based on a combination of template-based and ab initio docking followed by pyDock scoring [23**], which ranked 2nd and 1st among all the CAPRI predictors and scorers groups, respectively. Models for the subunits were built by CASP-hosted servers. Then, ab initio docking was applied in all cases, using appropriate symmetry constraints or interface restraints from literature. Additionally, when reliable templates were found, template-based models were built by superimposing all possible models of the monomers onto them. After sorting all built models by pyDock scoring, the proportion of template-based and ab initio docking models in the final set of submitted models depended on the reliability of the templates (Figure 2). The difference with other methodologies was more evident on the ‘difficult’ cases for which no clear template was available. For instance, in T154 ab initio docking by pyDock produced the only acceptable models among all participants. In T157, pyDock also produced some of the few successful models of all groups.

For scorers, pyDock was used to evaluate all the proposed models, and in case of reliable templates, consistency between energy-based scoring and template-based data was sought.

In 7th CAPRI, predictions using template information were in general successful. Indeed, failing to use available templates, as Fernández-Recio did in T122, T125 interface 1/4, and T133 targets, led to much worse predictions (although interestingly, this group was successful in the latter target, using only ab initio docking). This shows that it is critical to choose the optimal docking approach for each case, depending on the template availability. In the rest of targets, templates were used indirectly. In the two protein-peptide targets with good templates (T134, T135), ab initio docking with pyDock with restraints from the available templates was successful. In the six protein-saccharide targets (T126-130), ab initio docking on the cavity identified from the available templates was also successful. These represent alternative strategies to combine ab initio docking with template information.
Finally, in the scorers experiment, pyDock got the best performance when considering top 10 predictions, which shows its capabilities to evaluate complex models derived from combined approaches (template-based, ab initio, refinement) [24*] (Figure 1).

Novel methodological developments in protein docking

The most successful approach as predictor in CASP13-CAPRI was that of Venclovas group. They basically used template-based models when reliable templates were found, and free docking with HEX [11] otherwise. One of the reasons of their success could be the use of VoroMQA [28] for the evaluation and selection of the final models. However, they were less efficient in the scorers experiment (rank 7th), which might indicate that this function seems mostly optimized for their own pipeline for template-based and docking generation, while its application to models generated by other sources represents a challenge to be solved. Other successful approach was the use of CONSrank [29,30] for the ranking of docking models. CONSrank is based on the most frequent inter-residue contacts in the ensemble of decoys, and has been updated to Clust-CONSrank with the addition of a recently developed clustering procedure [31]. The best-performing server in CASP13-CAPRI was HDOCK [32], from Huang’s group, who developed a new pairwise shape-based scoring function (LSC) for protein–protein docking to take into account long-range interactions between protein atoms [33*].

Other recent new developments in protein docking are RosettaDock 4.0, which shows improved predictions for flexible cases [34*], LightDock, using glowworm swarm optimization with NMA-based flexible search [35], or CIPS, a new scoring procedure [36*] based on interface propensities from docking calculations. Docking interface propensities have interesting applications, such as interface prediction [37], and more recently, characterization

An example of the combination of template-based, ab initio docking and external data for integrative modeling of complexes. The scheme is based on the strategy followed by our group (Fernandez-Recio) as predictors in the recent CASP13-CAPRI and 7th CAPRI experiments.
of multi-protein complexes in combination with other evolutionary and physico-chemical properties [38].

Use of external information for integrative docking

The identification of correct docking poses often fails due to intrinsic errors in current scoring functions, incorrect consideration of oligomerization states, or because of multiple interfaces that are not usually included in docking calculations. For all these reasons, the use of external information on a given complex is often critical for successful docking predictions. The pioneering HADDOCK [16], as well as other protein–protein docking methods, such as pyDock [39], ZDOCK [40] or LightDock [4] have developed procedures to include distance restraints to improve the docking calculations. In this line, evolutionary information can be a relevant source of information for docking [42]. Indeed, the most successful docking approach in the recent 7th CAPRI edition was that of the Andreani and Guerois group. The challenging cases of this CAPRI edition encouraged them to go beyond their traditional rigid-body and InterEvScore approach, so they applied different strategies for the inclusion of evolutionary constraints, such as template-based modeling with RosettaCM-based protocol [43], identification of conserved anchoring interface motifs when only remote homologs were available, and covariance-based modeling of interacting subunits in cases in which traditional homology-based modeling would fail [44].

In a broader sense, integrative computational approaches that aim to efficiently use experimental structural data and additional information from a variety of sources for the structural modeling of complexes are becoming increasingly popular [45]. One example is the integration of Small-Angle X-ray Scattering (SAXS) experimental data in ab initio docking methods such as pyDock [46–48], HADDOCK [49], PatchDock [50,51], ATTRACT [52] or ClusPro [53]. And chemical cross-linking data has also been integrated in protein docking methods such as ZDOCK [54]. In the 7th CAPRI experiment, the use of integrative modeling approaches was blindly evaluated. Targets T150 and T151 were the same complex as T149, a challenging multi-domain dimer, for which SAXS and chemical cross-linking data were provided, respectively. Interestingly, the inclusion of restraints from SAXS data improved the models submitted by pyDock for the original target (with few successful groups), and the cross-linking data further improved pyDock submissions [55].

Conclusions

The most recent community-wide blind tests on the structural prediction of multi-molecular assemblies and heteromeric protein complexes (including interaction with peptides and saccharides) clearly showed that template availability, as well as any additional information on the complex, are critical for the modeling success. Several groups are focusing their efforts on developing new procedures for efficient integration of template-based and evolutionary information with ab initio docking methods, which are producing more accurate and realistic models. Additional methodological developments on protein docking include improvement of scoring functions, and better treatment of conformational flexibility during docking search, but the field is clearly moving towards an integrative analysis and modeling of protein complexes.

Conflict of interest statement

Nothing declared.

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