HawkDock: a web server to predict and analyze the protein–protein complex based on computational docking and MM/GBSA

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ABSTRACT

Protein–protein interactions (PPIs) play an important role in the different functions of cells, but accurate prediction of the three-dimensional structures for PPIs is still a notoriously difficult task. In this study, HawkDock, a free and open accessed web server, was developed to predict and analyze the structures of PPIs. In the HawkDock server, the ATTRACT docking algorithm, the HawkRank scoring function developed in our group and the MM/GBSA free energy decomposition analysis were seamlessly integrated into a multi-functional platform. The structures of PPIs were predicted by combining the ATTRACT docking and the HawkRank re-scoring, and the key residues for PPIs were highlighted by the MM/GBSA free energy decomposition. The molecular visualization was supported by 3Dmol.js. For the structural modeling of PPIs, HawkDock could achieve a better performance than ZDOCK 3.0.2 in the benchmark testing. For the prediction of key residues, the important residues that play an essential role in PPIs could be identified in the top 10 residues for ∼81.4% predicted models and ∼95.4% crystal structures in the benchmark dataset. To sum up, the HawkDock server is a powerful tool to predict the binding structures and identify the key residues of PPIs. The HawkDock server is accessible free of charge at http://cadd.zju.edu.cn/hawkdock/.

INTRODUCTION

Protein–protein interactions (PPIs) are involved virtually in all cellular processes, such as signal transduction, protein expression regulation and DNA replication. Therefore, determination of their complex structures is critical to understand the underlying molecular mechanisms of crucial biological processes (1) and even design compounds that interfere with PPIs with pharmaceutical significance (2). However, only a tiny number of the 3D structures of protein–protein complexes have been determined experimentally and deposited into the released databases, such as Protein Data Bank (PDB). In this context, computational methods, especially the protein–protein docking, have been increasingly applied to the structural prediction of macromolecular assemblies, which is expected to be a valuable complement to the experimental methods.

Protein–protein docking algorithms could be roughly classified into two categories: template-based modeling and template-free docking (3). Template-based modeling is based on the observation that protein–protein complexes usually interact in the same way if their interacting pairs share >30% sequence identity (4). Hence, the near-native structure of a protein–protein complex can be predicted by this method if an appropriate template is available. However, the number of the available templates is still very limited at present, and thus the template-free docking is generally more popular. Most template-free docking algorithms consist of two stages: sampling stage and scoring stage. In the sampling stage, a large number of decoys are generated, and the decoys sampled from this stage are re-scored and ranked by various scoring functions in the subsequent scoring stage.

A number of template-free docking servers have been developed and released to the public, such as ATTRACT (5), ClusPro (6), HADDOCK (7), ZDOCK server (8), SwarmDock (9), pyDockSAXS (10), pyDockWeb (11), InterEvDock2 (12), GRAMM-X (13), RosettaDock server (14), PatchDock (15), Hex server (16), 3D-Garden (17), FRODOCK 2.0 (18), etc. Moreover, a hybrid strategy by combining template-based modeling and template-free docking was proposed in HDOCK (19). The scoring strategies used in most servers rely on the scoring functions with a few energy terms, such as van der Waals potential, electrostatic potential, hydrogen-binding potential, desolvation,
etc. However, the desolvation energy, which plays a leading role in identifying correct binding poses, cannot be calculated accurately till now. In addition, most docking servers do not provide functions to identify and visualize the key residues for PPIs. It is quite challenging for non-expert users to identify the key residues in a protein–protein binding interface directly using some molecular visualization programs, as the binding interface usually involves a large number of residues (20).

In the past decade, two end-point free energy calculation methodologies, Molecular Mechanics/Poisson Boltzmann Surface Area (MM/PBSA) and Molecular Mechanics/Generalized Born Surface Area (MM/GBSA) (21), which are more theoretically rigorous than scoring functions, have been widely used to predict binding free energies and identify correct binding conformations for protein–protein systems (22–30). Furthermore, MM/PBSA and MM/GBSA with per-residue energy decomposition of the binding free energy of a protein–protein complex have been successfully employed to highlight the key residues in the binding interface (31–33). According to the assessment results reported by our previous study (25), MM/GBSA achieved better results not only in predicting the binding affinities but also in recognizing the near-native binding modes for PPIs than the tested scoring functions. However, MM/GBSA is much more computationally expensive than those commonly applied scoring functions due to the high computational cost of the polar desolvation term based on a Generalized Born (GB) model. In order to achieve a balance between computational efficiency and accuracy, we developed HawkRank, a physical scoring function with similar energy terms in MM/GBSA (34) by introducing a fast and effective method to calculate the desolvation potentials based on solvent accessible surface areas (35).

In this study, the HawkDock server was developed to predict and analyze the structures of PPIs by integrating the ATTRACT docking algorithm, the HawkRank scoring function and the MM/GBSA free energy decomposition analysis. In HawkDock, to predict the structure of a protein–protein complex, a large number of binding poses are first generated using rigid-body docking protocol of ATTRACT. Then, the near-native binding poses are recognized by the HawkRank scoring function combined with ATTRACT score. In addition, MM/GBSA was integrated into the HawkDock server to help users to analyze the key residues in a protein–protein binding interface and re-rank the top 10 models. All prediction and analysis functions in the HawkDock server are automated and the results are presented interactively through a user-friendly interface.

The MM/GBSA free energy decomposition

In our previous study, the MM/GBSA calculation based on the ff02 force field (39) and the GB{\textsuperscript{OB1}} model (40) yielded the best predictions for protein–protein binding free energies. Therefore, these parameters were used in the analysis of the key residues in protein–protein binding interfaces. First, all missing hydrogens and heavy atoms of the protein–protein complex were added by the \textit{leap} module in Amber16 (41), and then the force field parameters of the ff02 force field were assigned to the proteins. Subsequently, the complex was optimized in \emph{vacuo} by 2000 cycles of steepest descent and 3000 cycles of conjugate gradient minimizations. At last, the polar desolation energy was calculated by the modified GB (GB{\textsuperscript{OB1}}) models developed by Onufriev et al. The exterior and interior/solute dielectric constants were set to 80 and 1, respectively.

Input

The functions of the HawkDock server include the structural modeling of a protein–protein complex by using HawkDock and the prediction of the key residues in protein–protein binding interfaces using MM/GBSA.

Figure 2A shows the usage interfaces of HawkDock. The users are requested to upload PDB files or provide PDB
Figure 1. Workflow of the HawkDock server that is divided into three major steps: (i) input of unbound or bound protein structures; (ii) structural prediction of protein–protein complex by using the global docking algorithm implemented in ATTRACT and the HawkRank scoring function; (iii) identification of the key residues by MM/GBSA.

ID: chain ID (e.g. 1C3D:A). The non-standard amino acid residues, heteroatoms (HETATM records), original hydrogen atoms and residues with incomplete backbone atoms are removed and the missing side chains are added by PDB2PQR (42). In addition, the options of distance restraints and 're-rank top 10 models by MM/GBSA' are also available for the users. A number of binding poses will be generated in the local domain if distance restraints are provided. When the MM/GBSA re-scoring function is chosen, our server will not only re-rank the top 10 models predicted by ATTRACT and HawkRank, but also provide the top 10 most frequently occurring residues for the receptor and ligand found in the top 10 models. As the server runs, the job status information is displayed on the log page.

Figure 2B shows the usage interfaces of MM/GBSA. PDB files or PDB ID (e.g. 1SYX) should be input by the users. Similar to HawkDock, the non-standard amino acid residues, heteroatoms (HETATM records) and original hydrogen atoms are removed. The uploaded protein–protein complex structures need to be docked in advance or be de-
termed by experimental techniques. Moreover, the users are supposed to specify the chain IDs of receptor (e.g. A or A;B) and ligand (e.g. C or C;D). And the job status information of MM/GBSA is also displayed on the log page.

**Benchmarks**

The benchmark dataset I (Dataset I) to assess the structural modeling capability of HawkDock was obtained from ZDOCK benchmark 4.0 (43). ZDOCK benchmark 4.0 consists of 176 nonredundant protein–protein complexes, for which the NMR or X-ray unbound structures of the constituent proteins are also available, including the 124 cases in Benchmark 3.0 and 52 newly-added complexes. Since the 124 complexes in Benchmark 3.0 were used as the training set to develop HawkRank, the other 52 complexes were used as the benchmark dataset to evaluate the docking performance of HawkDock (Supplementary Table S1).

After carefully retrieving literatures, the benchmark dataset II (Dataset II) with 116 key residues in the complex interfaces identified by experiments for 43 proteins (receptors or ligands) in 32 complexes (a complex is separated into a receptor and a ligand) in Dataset I (Supplementary Table S2) was constructed to validate the prediction capability of MM/GBSA to determine the key residues for PPIs. The crystal structures were directly submitted to the MM/GBSA calculations and the key residues were then determined. As for a docking model, the binding pose with the smallest interface RMSD (I_RMSD) in the 10 000 decoys generated by HawkDock was analyzed by MM/GBSA.

**Evaluation criteria**

Two major evaluation parameters in the CAPRI campaign, ligand RMSD (L_RMSD) and I_RMSD, were used as the criteria to evaluate the docking performance for Dataset I. When I_RMSD is <4 Å or L_RMSD is <10 Å, the prediction is considered to be successful (44). The success rate was defined as the percentage of the total cases with at least one correct model in the top $N$ predictions.

For Dataset II, the success rate, defined as the percentage of the cases with at least one key residue within certain predictions, was employed to evaluate the key residue prediction capability of MM/GBSA.

**RESULTS**

**HawkDock server**

The HawkDock server based on the Python web framework of Tornado (an asynchronous networking library) is deployed on a Linux server of an Intel(R) Xeon(R) E5-2696 v3 2.30GHz CPUs with 36 cores and 64 GB of memory. After submitting jobs, the HawkDock server will create a HawkDock or MM/GBSA task and then put it into
the queue immediately. A HawkDock job usually needs 3 min (for proteins around 150 residues) to 25 min (for proteins ∼1000 residues). When the option of ‘re-rank top 10 by MM/GBSA’ is chosen, the running time of HawkDock will be extended to 25 min (for proteins ∼200 residues) to 175 min (for proteins ∼1000 residues). For MM/GBSA, the running time for the prediction of key residues is about 2 min (for proteins ∼200 residues) to 15 min (for proteins ∼1000 residues). Afterwards, the web interface will be redirected to the job status and result page. The URL of the page is unique for each job, and hence bookmarking the URL is highly recommended to the users. Moreover, an email will be sent to the user once the job is finished, if a valid email address has been provided by the user.

Output

Once the calculation is finished, the job status page will be auto-refreshed to the result page. As shown in Figure 3A, the result of HawkDock contains seven components. The downloadable files include three types of files: (i) the receptor and ligand PDB files used as the docking input; (ii) a text file with the docking scores for the top 100 models; (iii) the compressed tar files for the top 10 and top 100 models, respectively. If necessary, the users can also contact us for downloading more models.

In addition, the top 10 models can be interactively displayed with 3Dmol.js, a WebGL-based molecular viewer. The brief instructions of 3Dmol.js are shown on the bottom of the result page. It should be noted that all residues will be renumbered from 1 and their numbers might be different from those in the original file. Furthermore, the option is offered in the HawkDock result page to facilitate the users to directly perform the MM/GBSA analysis for one or several docking models predicted by ATTRACT and HawkRank. In the sixth part, the models can be selected and submitted directly to perform the MM/GBSA calculations. Moreover, the result page also shows the job name and the docking scores of the top 10 models.

If the users choose the option of ‘Re-rank top10 models by MM/GBSA’, as shown in Supplementary Figure S1, the top 10 most frequently occurring residues identified in the top 10 models for the receptor and ligand are displayed in the third and fourth parts, respectively. In addition, the analysis results for each model are listed in the eighth part. The top 10 models are re-ranked based on the binding free energies predicted by MM/GBSA.

The result page of MM/GBSA is presented in Figure 3B, including the job name, the value of the binding free energy of each complex, the downloadable file, the table sorted by the per-residue free energy contributions from the largest to the smallest, molecular viewer and its brief instruction. The upper and lower tables show the results of the receptor and ligand, respectively. The detailed values of all energy terms are summarized into a csv file. In addition, the users can highlight important residues in 3Dmol.js by selecting these residues in the table.

Docking performance on CAPRI targets

We participated in recent CASP13-CAPRI as a server group. According to the rules of CAPRI, the server groups must submit their results within 2 days, but the human predictor groups have ∼20 days and can use any additional information (6). Furthermore, only the sequences were provided for the docked systems so that the wrong monomer structure of proteins may be predicted from the beginning. Since it is the first time for us to participate in this challenge, we met some problems in the registration as a server group so that we missed some targets. In addition, HawkDock was designed for dimer docking and thus we ended up with participating in seven targets (T148, T152, T153, T154, T155, T156 and T157). As for the seven proteins in CASP13, taking the top 5 hits as the criterion, HawkDock could provide acceptable models for two targets (T152 and T153), compared with acceptable models for one target (T152) provided by LZERD (45) and acceptable models for two targets (T152 and T153) provided by other servers, such as ClusPro (6), SwarmDock (9), HADDOCK (7), HDOCK (19) and GalaxyPPDock (http://galaxy.seoklab.org/cgi-bin/submit.cgi?type=PPDock). Even for the human predictor groups, the best-performed groups could only give successful predictions for three targets. When the top 10 models were considered in the server groups, HDOCK and GalaxyPPDock performed better and achieved success for three targets. These results suggest the robustness of HawkDock in protein–protein docking.

Docking performance on benchmark

The results of HawkDock were also compared with those of ATTRACT and ZDOCK 3.0.2, a widely used rigid-body docking program (46), based on the analysis of Dataset I. 10 000 and 54 000 decoys were generated and ranked by ATTRACT and ZDOCK 3.0.2, respectively. The detailed ranking results are summarized in Supplementary Table S1 and the success rates from top 10 to top 1000 and top 1 to top 10 are shown in the main figure and the inserted zoomed view of Figure 4A, respectively. In terms of the top 10 level, HawkDock achieves a better performance with a success rate of 25.00%, while the success rates for ATTRACT and ZDOCK 3.0.2 are 13.46% and 21.15%, respectively. With more predictions considered, HawkDock performs significantly better than ATTRACT and ZDOCK 3.0.2 and yields the success rates of 42.31%, 50.00%, 69.23%, 80.77% and 88.46% at the top 50, 100, 200, 400 and 1000 levels, compared with 34.62%, 44.23%, 55.77%, 65.39% and 78.85% for ATTRACT and 30.77%, 42.31%, 50.00%, 65.39% and 71.15% for ZDOCK 3.0.2. At the top 1 level, HawkDock performs worse than ZDOCK 3.0.2, but its success rate can be improved from 5.76% to 11.54% by the MM/GBSA rescoring, which is similar to that of ZDOCK 3.0.2 and much higher than those of the other tested methods. Furthermore, the frequently occurring residues in the top 10 models can also be analyzed by this MM-GBSA refinement protocol. In summary, these results highlight the robustness and reliability of HawkDock in the structural prediction of PPIs.

Key residue prediction

The prediction capability of MM/GBSA in identifying the key residues in complex interfaces was then assessed based
Figure 3. The result pages of HawkDock and MM/GBSA. (A) At the top left of the page is (1) the job name, and under it are (2) the files for downloading. (3) A summary of the docking scores for the top 10 models is presented on the left bottom. (4) The top 10 models can be viewed in 3Dmol.js with (5) the optional buttons to control which model to display. (6) The table summarizes the submission and results of MM/GBSA and (7) the brief instructions are shown on the bottom. (B) At the top of page is (1) the job name, and under it is (2) the value of the binding free energy and (3) the files for downloading. The per-residue free energy contributions ordered from largest to smallest for (4) receptor and (5) ligand are displayed in the table, (6) through which the users can choose which residue to display in 3Dmol.js. (7) The brief instructions are also displayed on the bottom.
Figure 4. (A) The success rates from top 10 to top 1000 and top 1 to top 10 of HawkDock, ATTRACT, and ZDOCK 3.0.2, and (B) the success rates of the key residues predicted by MM/GBSA.

on the analysis of Dataset II. As expected, MM/GBSA showed an excellent performance in the key residue prediction (Supplementary Table S2 and Figure 4B). For the crystal structures in Dataset II, 39.53% (17) were successfully predicted at the top 1 level. Even for the predicted structures, MM/GBSA also reached a success rate of 23.26% at the top 1 level. As more predictions were considered in the evaluation, the success rates for both structures increased rapidly and were promoted to 95.35% and 81.40% in the top 10 level, respectively. Therefore, if a reliable model can be provided, an excellent performance can be achieved by MM/GBSA. In addition, analysis of the frequently occurring residues in the top 10 models were also executed by us. In dataset II, in terms of the top 10 level, the correct conformations can only be found in five proteins, four of which were predicted successfully at the top 5 level. For the proteins without an acceptable model at the top 10 level, 9 and 10 of 38 is also predicted successfully at the top 3 and 10 level, respectively, which provides an available solution when no reliable model is submitted. To sum up, MM/GBSA achieve a good performance in identifying the key residues in complex interfaces.

Examples of web server output

To illustrate the practicability of the HawkDock server, the complex 3D5S (47) from ZDOCK benchmark 4.0 predicted by the unbound structures of complement C3d fragment (C3d, PDB ID: 1C3D) (48) and fibrinogen-binding protein C-ter domain (Efb-C, PDB ID: 2GOM) (49) was used as an example. As shown in Figure 5, HawkDock generated the correct binding conformation with an 1RMSD of 1.337 Å between the crystal structure and the best hit. Then, this model was analyzed by MM/GBSA. The key residues of R131 and N138 from Efb-C (47) were successfully predicted in the top 10 predictions, ranking 4 and 10, respectively. Interestingly, the statistics of the frequently occurring residues in the top 10 models performed better than that based on the reliable model. R131 and N138 were successfully predicted at the one and eight rankings, respectively. For the crystal structure, it is clear to see that the MM/GBSA achieved an excellent performance and the key residue of N138 was successfully predicted as the best prediction (R131 was mutated to alanine in 3D5S).

CONCLUSION

Here, we present a user-friendly HawkDock server for the structural prediction and key residue analysis of PPIs. The combination of ATTRACT, HawkRank and MM/GBSA makes it very efficient for the identification of near-native docking models and key residues. The HawkDock server is an on-going project and further developments will be focus on the incorporation of additional information (e.g. co-evolutionary information or SAXS) and the integration with automatic modeling of protein structures.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.
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