idTarget: a web server for identifying protein targets of small chemical molecules with robust scoring functions and a divide-and-conquer docking approach

Jui-Chih Wang¹, Pei-Ying Chu², Chung-Ming Chen¹ and Jung-Hsin Lin²,³,⁴,*

¹Institute of Biomedical Engineering, National Taiwan University, ²Division of Mechanics, Research Center for Applied Sciences, Academia Sinica, ³School of Pharmacy, National Taiwan University and ⁴Institute of Biomedical Science, Academia Sinica, Taipei, Taiwan

Received March 4, 2012; Revised April 29, 2012; Accepted May 5, 2012

ABSTRACT

Identification of possible protein targets of small chemical molecules is an important step for unravelling their underlying causes of actions at the molecular level. To this end, we construct a web server, idTarget, which can predict possible binding targets of a small chemical molecule via a divide-and-conquer docking approach, in combination with our recently developed scoring functions based on robust regression analysis and quantum chemical charge models. Affinity profiles of the protein targets are used to provide the confidence levels of prediction. The divide-and-conquer docking approach uses adaptively constructed small overlapping grids to constrain the searching space, thereby achieving better docking efficiency. Unlike previous approaches that screen against a specific class of targets or a limited number of targets, idTarget screen against nearly all protein structures deposited in the Protein Data Bank (PDB). We show that idTarget is able to reproduce known off-targets of drugs or drug-like compounds, and the suggested new targets could be prioritized for further investigation. idTarget is freely available as a web-based server at http://idtarget.rcas.sinica.edu.tw.

INTRODUCTION

Identification of targets of small chemical molecules is essential for unravelling the underlying molecular causes of actions. Often, natural products, i.e. compounds discovered from plants, animals, marine lives or other living organism, exhibit useful pharmaceutical effects, e.g. anti-inflammatory, anti-cancer and anti-viral effects, yet their molecular mechanisms remain elusive. On the other hand, many drugs are known to be accompanied with unpleasant adverse effects, but the molecular targets of such effects are largely unknown. On the contrary, there are also some old drugs whose additional beneficiary effects are discovered only recently. For example, the epigenetic mechanism of the anticancer effect of cholesterol-lowering drugs, statins, was uncovered rather recently (1).

Conventional virtual screening of chemical libraries has been used widely to search for new leads in drug development for a protein target (2). As the deposited structures of biomolecules in the Protein Data Bank (PDB) increase substantially in the past decades, searching for the targets of a given drug or small compounds (also known as inverse screening, target fishing, off-target prediction, etc.) has become a useful approach (3–7).

One of the major hurdles for target identification is the effectiveness of scoring functions (7,8). To evaluate the binding affinity of the small ligand and a protein target, an accurate yet generally applicable scoring function is essential. We recently developed three robust scoring functions, AutoDock4 RRP, AutoDock4 RAP and AutoDock4RGG (9) based on the energetic terms and the formulation of AutoDock4 (10). These scoring functions report the binding free energy in the experimental scale, which allows direct comparison among different protein–ligand systems. Two of these three robust scoring functions were constructed using atomic charges from quantum chemical calculations, namely, RESP (11) and AM1-BCC (12), and the robust regression analysis (13) was employed to mitigate the influence of outliers for the calibration of the scoring functions. These robust AutoDock4 scoring functions have been benchmarked for their capability in binding affinity prediction and binding pose prediction (9). For the assessment of binding affinity prediction with a large external set of...
1427 complexes from PDBbind v2009, AutoDock4RAPP obtained root-mean-square errors of 2.176 kcal/mol, while the size of the training set is only 147.Benchmarked by using two decoy sets (14,15), the robust AutoDock4 scoring functions outperformed most of other scoring functions for the binding pose prediction (9).

Here, we utilize an efficient docking approach to screen the protein targets. Evaluation of potential targets is carried out by using the AutoDock4 robust scoring functions and the affinity profile analysis to enhance the confidence level of prediction.

**MATERIALS AND METHODS**

**Docking and scoring**

The search engine of idTarget web server is MEDock (16), which generates initial docking poses of the small ligand. The global search algorithm used in MEDock has also been tested recently by random mathematical functions simulating rugged free energy landscapes with different dimensionalities (17). It was shown that this global search algorithm maintained very high searching efficiency even at the dimensionality of 30 (17), which should be sufficient for applying to most protein–ligand systems. It was also shown that the traditional genetic algorithm failed to deliver good searching efficiency as long as the dimensionality is beyond 20 (17).

Since the binding sites are generally not known, the search space (i.e. the docking grid), should cover the entire protein. However, the searching efficiency also depends on the size of the search space. To achieve better searching efficiency, as well as to overcome the memory limitation when docking to very large proteins, a divide-and-conquer docking approach, similar to the partial box approach in BDT (18), was adopted for efficient blind docking. It should be emphasized that the entire receptor surface is searched, instead of only a part of the surface or just the active site. This also allows the search for possible allosteric binding sites. The divide-and-conquer docking approach can easily take the advantage of the parallel computing facilities and reduces the searching time by limiting the size of grids. The large box that covers the entire protein surface is subsequently divided into smaller boxes, where the size of the smaller boxes is dynamically determined according to the size of query ligand. If a grid box is far away from the receptor, namely, no atom is within 1.42 times the length of the grid box to the center, it will be eliminated to reduce the computational cost. The pre-calculated energetic grid maps for the protein targets are stored in the idTarget web server, and therefore the time for grid construction can be saved.

To analyze the results of target screening, the affinity profile of each binding site is constructed. Our affinity profile analysis was inspired by the structure-based maximal affinity model recently proposed (19). Here, instead of maximum affinity, the affinity profile of a given pocket is modelled by a Gaussian function, whose width is determined by the range of the predicted binding affinities of the complexes of different ligands in the same binding pocket of the same protein. A Z-score of ligand $j$ to a given protein pocket $i$ is calculated based on above profile using the following equation:

$$Z_{ij} = \frac{E_{ij} - E_i}{sd_i}$$

$$E_i = \sum_{k=1}^{N_c} \frac{E_k}{N_c}$$

$$sd_i = \sqrt{\frac{\sum_{k=1}^{N_c} (E_k^2 - E_i^2)^2 / N_c - 1}{N_c}}$$

where $E_{ij}$ is the dock energy of ligand $j$ docking to the protein pocket $i$. $E^c$ represents the energy directly evaluated with the crystal pose. $E_i$ and $sd_i$ are the center and the width of the affinity profile of protein pocket $i$, respectively. A large negative $Z$ value signifies an important target of the query compound.

**The contraction-and-expansion strategy**

Since direct screening for a large set of targets is computational intensive, a contraction-and-expansion strategy is adopted to perform target screening efficiently. In the contraction stage, the set of representative pockets for the protein–ligand complexes in the PDB, 3046 mean points of scPDB (20,21), were clustered by CD-Hit (22) under the cutoff of 40% sequence identity. After clustering of protein sequences in the PDB, a contracted list of targets (2091 targets) was obtained. The targets screening starts from this contracted list, subsequently, half of targets with lower docking energy are kept. In the expansion stage, targets that are homologous or contain similar binding sites to the top representatives are also selected for screening. The idTarget server is periodically updated to include new protein structures from PDB. The pockets of a new PDB entry will be compared with the set of representative pockets by SiteEngine (23) and subsequently assigned to the most similar pocket group.

In addition to the contraction-and-expansion strategy, idTarget provides two modes for searching binding poses. In the 'scanning mode', conventional docking jobs are performed for each protein structure. In the 'fast mode', the ligand will first be mapped to the binding sites of the homologous proteins by superposition of homologous protein structures with the pre-calculated structural alignment information by using CE (24), and then its binding poses are optimized by adaptive local searches to remove too close contacts with protein atoms. To extend the coverage of PDB structures, idTarget also performs a ligand similarity search to find structurally similar co-crystallized ligands. The Tanimoto coefficients calculated by OpenBabel 2.3.0 (25) are used to rank the results of the similarity search by matching molecular fingerprints.

**INPUT**

The input window of the idTarget web server requires an input ligand file for the target screening. Users can upload a ligand file with the popular molecular file formats (pdb,
three examples are presented here. The first example is
To demonstrate the usage of the idTarget web server, EXAMPLES web/).
OpenAstexViewer applet (http://openastexviewer.net/ resulting 3D structures interactively via the embedded can inspect the binding poses and manipulate the target name are collected in the third layer. Users can also query a small set of proteins with known structures by editing a PDB ID list.

OUTPUT
Tracking the status of a job and retrieving the results can be achieved by input the Task ID. If the e-mail address is provided, the notification will be sent to users as soon as the calculation is finished. Although it usually takes hours to finish a job, depending on the loading of the server, users can always check the results while some targets have been screened. During the calculation of a job, the screening results are sorted immediately for showing the potential targets in the higher ranks. As depicted in Figure 1, two hierarchical tables and a molecular graphic visualizer are shown in the result page. The two-layer table (grouped by protein name) lists all the results sorted by dock energies. The PDB entries with the same target name will be grouped, and one can click on the button to expand/collapse the next layer or all results at once. The homologous proteins are grouped in the three-layer table (grouped by the binding pocket), where the representative one is shown in the first layer and the proteins within the same binding pocket are shown in the second layer. All the PDB entries with the same target name are collected in the third layer. Users can also use selected ranges of predicted binding free energy and Z-score as the filtering criteria to obtain a more condensed target list.

WEB SERVER
The web interface of idTarget was written in PHP (5.1.6). Marvin Sketch applet (http://www.chemaxon.com/products/marvin/marvinsketch/) was embedded in the idTarget job submission page so that users can draw their interested ligand online. In the result page, users can inspect the binding poses and manipulate the resulting 3D structures interactively via the embedded OpenAstexViewer applet (http://openastexviewer.net/web/).

EXAMPLES
To demonstrate the usage of the idTarget web server, three examples are presented here. The first example is
the HIV-1 protease inhibitor, darunavir (DRV), which has been approved by FDA on 2006 and has been validated the inhibition against multidrug-resistant viruses (26). The second example is one of indirubin derivatives, 6-bromo-indirubin-3’ oxime (6BIO), which has been experimentally validated as an inhibitor of several kinds of protein kinases (27). The third example is an inhibitor of histone deacetylase 2 (HDAC2), namely, N-(4-aminobiphenyl-3-yl)-benzamide (dubbed ‘LLX’ from its PDB component ID). The target screening results of these three examples are archived in the web server with the Task IDs DRV_screening, 6BIO_screening and LLX_screening, respectively.

DRV
Mutations of HIV-1 protease (PR) residues usually reduce susceptibility to the PR inhibitors but not significantly to darunavir. There have been many structural and kinetic analyses of darunavir against several drug-resistant mutants (26,28–30). Table 1 gives the comparison of predicted and experimental binding affinities for the wild-type and mutant HIV-1 PR. These results indicate the comparable predicted binding affinity to the experimental data. Although the scoring function was not trained with these PR-inhibitor complexes, it still can discriminate the fine difference of affinity against mutants, except for the case of 3CYW. It should be noted that in the docking, we have excluded a well-known flap water molecule, which forms a hydrogen bond network between the flaps of PR and ligand. If that flap water was included in docking of DRV, the predicted free energy will be lowered by ~1 kcal/mol. This flap water was removed because in general it may cause steric clashes for most ligands. Figure 2 depicts the binding pose of DRV against the wild-type PR with the rmsd of 0.85Å compared to the crystal pose.

6BIO
This example shows that not only the well-known kinase targets (CDK2, CDK5 and GSK-3β) could be identified by idTarget, but also the kinase PDK1 (27) and other proteins. More detailed processes are described in ‘Example 1’ in our idTarget web server. The predicted best targets are as follows: CDK2 (PDB ID: 1FVV, predicted $Ki$: 51.7 nM); CDK5 (PDB ID: 1UNH, predicted $Ki$: 71.2 nM); GSK-3β (PDB ID: 1Q41, predicted $Ki$: 63.3 nM); PDK1 (PDB ID: 2PE1, predicted $Ki$: 96.5 nM). Figure 3 depicts the best docking pose of 6-bromo-indirubin-3’ oxime (6BIO) in PDK1 that confirms the proposed pose in Zahler et al. (27).

To compare directly with the results of Zahler et al. (27), we performed the screening with the same set of 5821 PDB entries by using the ‘User-edited list’ feature. The predicted binding free energy was used to rank the screening results, and the Z-score (<0) was the filter. For the known protein kinase targets (CDK2, CDK5 and GSK-3β), the topp 1% and top 5% enrichment factors in our screening results were 6.45 and 11.94, respectively, while the top 1% and top 5% enrichment factors calculated from the results of Zahler et al. were 19.35.
and 7.42, respectively. It should be noted that protein kinases are the only targets of interest in the work of Zahler et al. However, we found that two proteins, matrix metalloproteinase and glycogen phosphorylase, constitute the major part of the top 1% of our results. Interestingly, we found that a recent experimental work (31) showed that an indirubin derivative inhibits rabbit muscle glycogen phosphorylase. The enrichment factor for glycogen phosphorylase (PYGM) in the top 1% was 18.18 in our results, but none of PYGM was identified in the top 5% of the results of Zahler et al. We further performed the ROC curve analysis (32) to assess the performance of the prediction. Figure 4 shows the ROC curves and numerical values of the areas under the curve (AUC), where CDK2, CDK5 and GSK-3\(\beta\) are considered as known targets of 6BIO (in Figure 4a), and these three kinases plus PYGM are considered as known targets (in Figure 4b), respectively. Supplementary Figures S1–S4

Table 1. DARUNAVIR screening results for wild-type and mutant HIV-1 PR.

<table>
<thead>
<tr>
<th>PDBID</th>
<th>(\Delta G^{\text{pred}}) (kcal/mol)</th>
<th>(K_i^{\text{pred}}) (nM)</th>
<th>Z-score</th>
<th>mutant</th>
<th>(K_i^{\text{exp}}) (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3CYW</td>
<td>-10.99</td>
<td>8.8</td>
<td>0.26</td>
<td>G48V</td>
<td>17a</td>
</tr>
<tr>
<td>2F81</td>
<td>-10.82</td>
<td>11.7</td>
<td>0.32</td>
<td>L90M</td>
<td>0.03b</td>
</tr>
<tr>
<td>2IEN</td>
<td>-10.79</td>
<td>12.3</td>
<td>0.33</td>
<td>wild</td>
<td>0.22-1.0g, h, c</td>
</tr>
<tr>
<td>2IDW</td>
<td>-10.65</td>
<td>15.6</td>
<td>0.38</td>
<td>V82A</td>
<td>0.8-1.3h, c</td>
</tr>
<tr>
<td>2IEO</td>
<td>-10.55</td>
<td>18.5</td>
<td>0.41</td>
<td>I84V</td>
<td>3.2h, c</td>
</tr>
<tr>
<td>1T3R</td>
<td>-10.53</td>
<td>19.1</td>
<td>0.42</td>
<td>wild</td>
<td>0.06d</td>
</tr>
<tr>
<td>2HSI</td>
<td>-10.4</td>
<td>21.1</td>
<td>0.44</td>
<td>V32I</td>
<td>3.3c</td>
</tr>
<tr>
<td>2F80</td>
<td>-10.4</td>
<td>23.8</td>
<td>0.46</td>
<td>D30N</td>
<td>6.0</td>
</tr>
<tr>
<td>2F8G</td>
<td>-10.38</td>
<td>24.6</td>
<td>0.47</td>
<td>I50V</td>
<td>2.0-18h</td>
</tr>
<tr>
<td>3D1Z</td>
<td>-10.38</td>
<td>24.6</td>
<td>0.47</td>
<td>I54M</td>
<td>1.6h</td>
</tr>
<tr>
<td>3D20</td>
<td>-10.38</td>
<td>24.6</td>
<td>0.47</td>
<td>I54V</td>
<td>5a</td>
</tr>
</tbody>
</table>

show the ROC curves with different selections of known targets and Z-score filters. The more the stringent of the Z-score filter, the higher the AUC was obtained, which could be due to the nature of conformational selectivity, the general applicability and the universal physical scale of our robust scoring function (9). These analyses also indicate that our predictions are comparable with the prediction of Zahler et al.

**LLX**

N-(4-aminobiphenyl-3-yl)-benzamide is a compound recently designed and synthesized to optimize the fitting of the HDAC2 binding pocket (33). It has been long recognized that there is a 14-Å long internal cavity, also called 'foot pocket', adjacent to the catalytic zinc ion of HDAC. This foot pocket connects to a lipophilic tube with an angle of about 110°, and this compound sits at the kink between the foot pocket and the tube. As shown in Figure 1, the screenshot of the result page given by idTarget, the compound LLX recognized its co-crystalized protein structure (PDB ID: 3MAX) with the highest affinity.

**DISCUSSION**

Although darunavir is a relatively new drug, its adverse effects and interactions with other drugs have been reported (34,35). It may be possible to further investigate possible molecular causes of actions of these adverse effects and drug–drug interactions by scrutinizing the list of proteins identified by idTarget. Our prediction for possible targets of 6BIO is generally consistent with the previous work (27). It is interesting to note that the top prediction of LLX is 3MAX, which is the HDAC2 structure crystallized by the same group that designs and synthesizes this N-(4-aminobiphenyl-3-yl)-benzamide. A possible explanation could be that this compound sits at the kink between the foot pocket and the lipophilic tube, which altogether forms a very curved channel structure and may be rather unique among protein structures.

Drugs can bind to a myriad of particles in the blood, such as red blood cell, leukocytes, platelets, as well as to some proteins, e.g. serum albumin, lipoproteins, α1-acid glycoproteins, α, β, γ-globulins, etc. (36). Usually, the
REFERENCES


