INTRODUCTION

Protein–ligand docking has been widely used to predict binding modes and affinities of ligands. Protein–ligand docking is a powerful tool for computer-aided drug discovery (CADD). Currently, there are dozens of commercial and academic tools available for protein–ligand docking [1-12]. Most docking tools require the ligand binding region (the rotation and translation of a ligand in this region) in advance to search for the most energy favorable binding mode. The binding region is usually represented as a cubic box, so its size and center are critical for accurate docking because it defines the boundaries of the conformational sampling space. In many application scenarios, the binding regions are unknown. To identify potential interactions between a given protein and a ligand, docking has to be performed on the entire protein surface to find the most probable binding mode. This process is called blind docking [13-16]. Compared to regular docking, blind docking is less reliable and stable as the docking space is usually too large to sufficiently sample using a limited number of random searches. Nevertheless, blind docking is particularly valuable for discovering unexpected interactions that may occur in unidentified binding modes [17].

Traditionally, blind docking is performed on the entire protein surface. Alternatively, docking on putative binding regions of the given protein usually improves the sampling efficiency and reduces the computational cost of blind docking [18]. Currently, many binding site detection tools have been developed [19-29]. These methods help users find residues that potentially bind with ligands. However, users must cluster residues into groups and estimate the parameters manually and then perform several rounds of protein–ligand docking to obtain the final result. Although this process is feasible, it is not efficient and has not been systematically optimized. To address this problem, several blind docking tools have been developed in recent years that have integrated cavity detection with a focused docking module. For example, popular software SwissDock [30, 31], QuickVina-W [15] and BSP-SLIM [32] provide particularly valuable services for blind docking. In this paper, we described a new blind docking tool, named CB-Dock, which focuses on enhancing the docking accuracy. CB-Dock predicts binding regions of a given protein, calculates the centers and sizes with a curvature-based cavity detection approach, and performs docking with a popular docking program, Autodock Vina. This method was carefully optimized and achieved ~70% success rate for the top-ranking poses whose root mean square deviation (RMSD) were within 2 Å from the X-ray pose, which outperformed the state-of-the-art blind docking tools in our benchmark tests. CB-Dock offers an interactive 3D visualization of results, and is freely available at http://cao.labshare.cn/cb-dock/.

Keywords: bioinformatics; computer-aided design; computer-aided drug discovery

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**MATERIALS AND METHODS**

**Benchmark dataset**

**PDBbind Set.** A total of 1684 protein–ligand structures were selected from PDBbind (v2018) [8, 34]. The molecular weights of the proteins were limited to 150–500 g/mol, and the numbers of rotatable bonds were within 10. In addition, the proteins that share 60% or more similarity to the Astex Diverse Set or MTiAutoDock data were eliminated. The structures can be downloaded from our website [http://cao.labshare.cn/cb-dock/](http://cao.labshare.cn/cb-dock/).

**Astex Diverse Set.** The Astex Diverse Set contains 85 protein–ligand complexes [35], which were downloaded from the Protein Data Bank [36]. The redundant identical chains, water molecules, and heteroatoms were discarded.

**MTiAutoDock Set.** The test data are from the benchmark set of MTiOpenScreen [37]. The data contains 27 crystal structures that cover important drug targets, including enzymes, GPCRs, nuclear receptors, and PPIs.

**RESULTS**

**Detecting cavities on proteins**

Most small-molecule binding occurs in protein pockets or cavities because high affinity can only be gained by sufficiently large interaction interfaces [40]. CB-Dock searches for concave surfaces to detect cavities. Briefly, CB-Dock generates a set of points to represent the solvent-accessible surface and calculates the curvature factor of each point using the method from our previous work [41, 42]. These points at the concave surface (curvature factor > 8) are clustered by a density-peak-based clustering algorithm [43]. Thus, we obtained several clusters of points that represent cavities on the protein surface. We present the example of aminopeptidase (PDB ID: 1TXR), whose cavities are highlighted in Fig. 1a. The cavities were ranked according to their sizes. We compared our method (called CurPocket) with state-of-the-art protein–ligand binding site prediction methods using the benchmark set of COACH [23], which is one of the best prediction methods. The results showed that our method is comparable to that of COACH in terms of Matthews correlation coefficient, precision, and recall (see Supplementary Table S1). Unlike traditional binding site prediction methods, our method detected the real binding cavities as much as possible to offer options for blind docking. To investigate its performance in detecting real binding cavities, we submitted 1684 structures from PDBbind to CurPocket (see the Materials and methods section) and examined their success rates by comparing the top 10 cavities with the real binding cavities from the crystal structures. Test results showed that the predicted success rates [44] of the top 1 to 10 cavities increased from 63.7% to 92.4%, respectively (Fig. 1b). From the top 10 to top 5, the success rate dropped only 2%. To balance the computational expense and cavity detection accuracy, we selected the top 5 cavities as candidates for focused docking.

**Calculating centers and sizes of docking boxes**

For a putative cavity, CB-Dock needs to customize a docking box for the following computation. A good docking box should enclose the native binding pose and exclude as many as possible irrelevant poses. The center and size of the docking box are the key parameters in this process. The center of the ligand from the crystal structure is the best choice for the docking box; however, we can base these parameters only on the putative cavity and its surrounding area.
The performances of these methods were quantified by the percentage of correct predictions (RMSD < 2 Å) (Fig. 3a). The results show that for traditional blind docking, redocking, and CB-Dock, the prediction accuracies were 38.8%, 39.5% and 69.4%, respectively. As we expected, CB-Dock had significant improvements (~30% higher) over traditional blind docking, and the overall accuracy was much closer to redocking and the upper limit of docking using Autodock Vina. Particularly, when the prediction was correct, CB-Dock and redocking had nearly identical RMSD values (Fig. 3b). This result implies that the cavity detection and docking parameters of CB-Dock work rather well. As Autodock Vina is based on a random algorithm, whose results may be different from the repeat runs, we repeated the test for 3 rounds to investigate the stability of the three methods. The results showed that the RMSD variations of CB-Dock and redocking were less than 5%, while it was up to 10% for traditional blind docking. We argued that CB-Dock appropriately decreased the sampling space and thereby reduced the randomness of the results. In all, cavity detection is a powerful approach to improve blind docking.

Comparison of CB-Dock with existing blind docking tools

To gain an overall performance of CB-Dock, we further compared it with four state-of-the-art docking tools, including DockingApp [38], MTIAutoDock [37], rDock [45], and SwissDock [30, 31]. Though the tools provide multiple usages, we focused on their performance of blind docking. DockingApp searches for binding sites over the whole protein surface by AutoDock Vina [33]. MTIAutoDock employs a variant of the grid-based LIGSITE algorithm [46] to identify cavities. In general, DockingApp and MTIAutoDock follow the traditional strategy, while rDock, SwissDock, and CB-Dock only allow docking in the putative binding regions. We conducted the benchmarks on the Astex Diverse Set and MTIAutoDock data (see the Materials and methods section). In the first dataset, DockingApp, MTIAutoDock, rDock, SwissDock (accurate mode) and CB-Dock achieved 42.4%, 42.4%, 41.2%, 53.0%, and 69.4% success rates of top-ranking poses within the RMSD of 2 Å from crystal structures (see the Materials and methods section). In the second set, the five tools achieved 33.3%, 51.9%, 33.3%, 70.4% and 74.1% success rates, respectively (Fig. 4b). Both benchmarks illustrated that, in terms of success rates for top-ranking poses, CB-Dock outperformed other blind docking tools. As blind docking strongly depends on the

unbound ligands to estimate the center and size. Hence, we first selected the center of the putative cavity, i.e., the center of points at the concave surface, as the docking center. To quantify its deviation from the best center, we calculated distances between the two centers using the PDBbind data set (see the Materials and methods section). The distances between centers of real and putative target cavities were distributed from 1 to 10 Å (Fig. 2a). For most of the data (76.6%), the distances were within 5 Å and up to 97.7% when distances were within 10 Å. The result indicated that for the majority of the data, the center of the cavity was close to the ideal center. Second, we needed to determine the lengths of the docking box in each dimension, which was related to the size of the cavity, the size of the ligand and the deviation of the putative center from the ideal center. After systematic examination of the outcome from docking, we finally calculated the i axis length $L_i$ of the docking box by a constant $x$ plus the maximum of the length $C_i$ of the putative cavity or gyration radius $R$ of the given ligand as follows:

$$L_i = x + \max(R, C_i)$$

The constant $x$ is used to compensate for the deviation of the putative center and to ensure that the ligand is enclosed in the docking box. To determine $x$, we tested the above protein–ligand structures to investigate the proportion of docking boxes that enclosed the ligands by gradually increasing $x$ from 0 to 12 Å (Fig. 2b). The results showed that the proportion grows rapidly when $x$ increases from 0 to 5 Å. When $x = 10$ Å, all the ligands are enclosed in the docking box. Thus, we choose $x = 10$ Å in our program. Detailed analysis shows that the sizes of the docking box by the above formula were mostly less than 30 Å, which was within the recommended upper limit (http://vina.scripps.edu/manual.html#faq).

The guidance of cavity detection improved blind docking

To assess the performance of CB-Dock, we compared it with traditional blind docking using a protein–ligand complex from Astex Diverse Set [35]. The docking parameters of traditional blind docking are described in section ‘The traditional blind docking and redocking protocols’. In addition, to determine the upper limit of this blind docking, we also tested redocking the centers and sizes of docking boxes that were obtained from crystal structures [39]. We measured the accuracy by RMSD between the predicted binding mode with the lowest docking score and the native mode in the crystal structures. The performances of these methods were quantified by the percentage of correct predictions (RMSD < 2 Å) (Fig. 3a). The results show that for traditional blind docking, redocking, and CB-Dock, the prediction accuracies were 38.8%, 39.5% and 69.4%, respectively. As we expected, CB-Dock had significant improvements (~30% higher) over traditional blind docking, and the overall accuracy was much closer to redocking and the upper limit of docking using Autodock Vina. Particularly, when the prediction was correct, CB-Dock and redocking had nearly identical RMSD values (Fig. 3b). This result implies that the cavity detection and docking parameters of CB-Dock work rather well. As Autodock Vina is based on a random algorithm, whose results may be different from the repeat runs, we repeated the test for 3 rounds to investigate the stability of the three methods. The results showed that the RMSD variations of CB-Dock and redocking were less than 5%, while it was up to 10% for traditional blind docking. We argued that CB-Dock appropriately decreased the sampling space and thereby reduced the randomness of the results. In all, cavity detection is a powerful approach to improve blind docking.

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The above tests benchmarked blind docking on the ligand-bound states (holo) of receptors from the protein–ligand complex structures. The blind docking in unbound (apo) structures is much more challenging as the conformational changes of proteins are difficult to predict. We performed blind docking using the 19 apo crystal structures available in Astex Diverse Set (see Apo Structure Set in the Materials and methods section). The results showed that the average percentages of correctly predicted binding sites [44] of the top-one predictions are 47.4%, 36.8%, 47.4%, 31.6%, and 68.4% for DockingApp, MTiAutoDock, rDock, SwissDock (accurate mode), and CB-Dock, respectively.
and CB-Dock, respectively. The RMSDs exhibited a similar trend. The success rates of top-ranking sites within the RMSD of 5 Å are 36.8%, 31.6%, 42.1%, 26.3%, and 63.2%, respectively (see Table 1). CB-Dock achieved the highest accuracy in the Apo Structure Set. However, the success rate was notably lower than that on the holo structure set. Analysis showed that the conformational differences between apo and holo structures may result in two types of inaccurate docking. One type is that CB-Dock identifies accurate cavities for docking; however, the detailed conformation of cavities was different between apo and holo structures. The differences were critical for binding, docking may not be accurate because CB-Dock does not model the conformational changes between apo and holo structures. An example of this type is the PDB structure 1L2S (see Fig. S1a and S1b). The side chain of Ser64 at the protein side to be in the PDB format and a ligand file in the MOL2, MOL, or SDF. After submission, CB-Dock checks the input files and converts them to pdbqt formatted files using OpenBabel [47] and MGLTools [5]. Next, CB-Dock predicts cavities of the protein and calculates the centers and sizes of the top N (n = 5 by default) cavities. Each center and size, as well as the pdbqt files, are submitted to AutoDock Vina for docking. The final results are displayed after the computation of N rounds. Users can browse binding scores, cavity sizes, and docking parameters of the predicted binding modes in a table. Moreover, users can inspect the 3D structures of any binding modes on the web page by clicking the structures in the related table. The interactive 3D structures are drawn by NGL Viewer [48], which is supported by most modern browsers. Users are able to display atom-specific information, rotate and translate molecules, select models and colors. For more details, users could refer to the manual on the CB-Dock homepage.

Here, we present a case study of the software CB-Dock (Fig. 5). Nutlin3a, a potential anti-cancer drug, is able to bind with the E3 ubiquitin-protein ligase MDM2 and inhibit the MDM2–P53 interaction. The MDM2 protein structure was downloaded (PDB ID: 4HG7) from PDB. The Nutlin-3a mol2 file was generated by the PRODRG software [49]. The two files were uploaded and

### Table 1. The RMSDs of five blind docking tools benchmarked in Apo Structure Set

| Target protein | DockingApp (Å) | MTiAutoDock (Å) | rDock (Å) | SwissDock (Å) | CB-Dock (Å) |
|----------------|----------------|-----------------|-----------|---------------|-------------|
| 1hq2           | 3.019          | 12.796          | 3.018     | 3.601         | 3.037       |
| 1ke5           | 7.543          | 21.849          | 16.685    | 37.472        | 7.535       |
| 1l2s           | 15.528         | 3.874           | 7.361     | 31.838        | 15.497      |
| 1l7f           | 26.68          | 19.626          | 34.651    | 17.328        | 0.865       |
| 1n1m           | 34.526         | 18.808          | 24.157    | 21.524        | 23.342      |
| 1n2v           | 22.755         | 3.903           | 3.836     | 27.137        | 4.517       |
| 1ogo           | 17.044         | 12.092          | 14.372    | 5.265         | 1.817       |
| 1oyt           | 0.339          | 0.538           | 4.766     | 3.783         | 0.335       |
| 1q41           | 2.219          | 1.382           | 29.771    | 1.483         | 2.145       |
| 1s3v           | 3.912          | 4.098           | 1.802     | 3.243         | 4.622       |
| 1t40           | 4.750          | 30.219          | 28.091    | 30.17         | 4.726       |
| 1t46           | 17.22          | 32.354          | 18.139    | 17.82         | 17.206      |
| 1v0p           | 37.477         | 4.673           | 36.866    | 23.273        | 4.212       |
| 1v48           | 7.971          | 15.968          | 13.423    | 13.615        | 7.908       |
| 1w1p           | 9.203          | 10.918          | 1.721     | 27.866        | 13.354      |
| 1yvf           | 16.433         | 60.384          | 19.995    | 13.263        | 14.687      |
| 1ywr           | 4.428          | 25.78           | 4.284     | 24.156        | 4.381       |
| 2br1           | 4.383          | 5.081           | 3.749     | 1.589         | 4.381       |
| 2bsm           | 16.412         | 16.78           | 3.744     | 20.386        | 0.784       |
| Average        | 13.255         | 15.849          | 14.233    | 17.095        | 7.124       |
| RMSD < 5 Å     | 36.8%          | 31.6%           | 42.1%     | 26.3%         | 63.2%       |

The RMSD values < 5 Å are highlighted in bold
submitted to the CB-Dock server by clicking the button “Submit”. While processing docking, a progress bar appeared to indicate the status of the job. When the processing was complete (after approximately 2 min), the web page was updated with the results. The table listed Vina scores, cavity sizes, docking centers, and sizes of predicted cavities. Once a ligand in the table is selected, the structure in the interactive 3D graphics is visualized. In our example, the top binding mode with a Vina score of $-8.4$ also had the largest binding cavity. The binding mode was almost identical to the mode of ligand in the crystal structure (RMSD = 0.484 Å).

**DISCUSSION**

Discovering protein–ligand binding sites and conformations are particularly important in drug discovery. Blind docking is a powerful method for obtaining that information. Blind docking is also one of the key components in high-throughput screening and inverse docking [50–53]. Therefore, it is of great value to develop accurate blind docking tools. Thanks to the well-established AutoDock Vina docking software, we focused on developing methods of cavity detection and docking parameter optimization, which are critical for blind docking. CB-Dock is the first cavity detection-guided blind docking tool designed with AutoDock Vina docking software, we focused on developing methods of cavity detection and docking parameter optimization, which are critical for blind docking. CB-Dock is the first cavity detection-guided blind docking tool designed with AutoDock Vina among many popular Vina-based tools (http://vina.scripps.edu/manual.html#faq). The benchmark tests show that CB-Dock outperforms other state-of-the-art blind docking tools in terms of predicting binding sites and binding conformations. This performance is attributed to the curvature-based cavity detection that precisely narrows down the docking space as well as the optimized parameters for AutoDock Vina. Some shortcomings of CB-Dock were also observed in the test. First, compared to regular docking, CB-Dock was more time expensive because the docking was performed iteratively in five cavities. To reduce time consumption, cavity detection should be further improved in the future. Second, if the size of cavities was notably greater than that of the ligand, the accuracy of docking tends to decrease. A typical example is the huge cavity detected on nitric-oxide synthase (PDB ID: 1MMV), in which the predicted docking position is at the opposite side of the cavity (see Fig. S3). This result is mainly related to the accuracy of the scoring function, which is supposed to distinguish the global minimum from local minimums. Using an additional scoring function to rerank binding positions could be a solution to this problem. Third, CB-Dock needs to improve the accuracy of docking in apo structures. Compared to holo structures, apo structures show conformational rearrangement in ligand binding sites, which has not been captured in current CB-Dock software. In the following developments of CB-Dock, the protein conformation sampling method will be incorporated in CB-Dock to enhance docking in apo structures.

Apart from blind docking capabilities, user-friendly interfaces are also very important for docking tools. CB-Dock offers a convenient web service that allows even nonexpert users to perform protein–ligand docking and visualize results in 3D. We believe that CB-Dock can contribute to the characterization of newly determined protein structures and suggest novel therapeutic targets for biological and pharmaceutical studies.

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**AUTHOR CONTRIBUTION**

YL designed and optimized the CB-Dock tool and wrote the manuscript. MG built the CB-Dock web server. WTD benchmarked the program. MCH tested the server. ZXX guided the experiments. YC designed the project and wrote the manuscript.
ADDITIONAL INFORMATION
The online version of this article (https://doi.org/10.1038/s41401-019-0228-6) contains supplementary material, which is available to authorized users.

Competing interests: The authors declare that they have no conflict of interest.

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