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MedusaScore: An Accurate Force-Field Based Scoring Function for Virtual Drug Screening

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Abstract

Virtual screening is becoming an important tool for drug discovery. However, the application of virtual screening has been limited by the lack of accurate scoring functions. Here, we present a novel scoring function, MedusaScore, for evaluating protein-ligand binding. MedusaScore is based on models of physical interactions that include van der Waals, solvation and hydrogen bonding energies. To ensure the best transferability of the scoring function, we do not use any protein-ligand experimental data for parameter training. We then test the MedusaScore for docking decoy recognition and binding affinity prediction and find superior performance compared to other widely used scoring functions. Statistical analysis indicates that one source of inaccuracy of MedusaScore may arise from the unaccounted entropic loss upon ligand binding, which suggests avenues of approach for further MedusaScore improvement.

Introduction

The current rate of emerging new pharmaceutical targets outpaces that of new drug leads ¹, posing a significant challenge for drug discovery. Traditional trial-and-error approach to drug discovery represents a substantial challenge due to the enormous dimensionality of the chemical space. Structure-based drug design is a promising approach of rational drug discovery, which takes advantage of the increasing amount of solved three-dimensional structures of target proteins². By computational modeling of the target binding site, a ligand can be constructed de novo³ or via screening over a large database of millions of chemical compounds (virtual screening)⁴, ⁵. In a typical virtual screening workflow, a library of small molecules are first computationally docked to the target protein, and then ranked according to the predicted binding affinity. A scoring function is used throughout the process to (a) recognize the correct binding pose out of hundreds of computer-generated docking models (decoys), and to (b) predict the binding affinity for each molecule.

Docking algorithms have undergone substantial developments over the last two decades⁴, 6-12. Early attempts treated receptors and ligands as rigid bodies and docking was only based on molecular surface matching¹³. Docking programs nowadays not only allow full flexibility of the ligands but also partially treat receptors as flexible objects. However, how to systematically treat protein flexibility and ligand-induced protein conformational changes remains a considerable challenge in the further development of docking algorithms⁴. Despite

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the progress in docking methodologies, recent studies have shown that scoring functions are becoming a bottleneck in structure-based virtual screening $^{14\text{-}16}$. Benchmark studies have found that in many cases, docking programs can generate native-like docking poses, but these are often missed at the scoring stage. Moreover, the binding scores predicted by the scoring functions may exhibit poor correlation with the actual binding affinity, resulting in a large percentage of false positives in the "hit list" 16 .

Although several types of traditional scoring functions exist, all of them have major limitations. Physical force field-based approaches aim to describe protein-ligand interactions using elementary physical interactions. Combined with molecular dynamics (MD) and free energy perturbation (FEP) techniques 17 , 18 , the binding affinity can be reproduced to within an accuracy of 1 kcal/mol 19 , 20 . However, such approaches often involve intensive computation, especially in connection with the explicit solvent used to describe desolvation effects and hydrogen bonding. Because of the sampling insufficiency, such approaches are often limited to structurally similar targets and ligands. These speed and sampling limitations undermine the application of the physical force field-based scoring functions in virtual screening of vast molecular libraries. Alternatively, empirical and knowledge-based scoring functions 7 , $^{21-27}$ circumvent the speed and sampling problems by dissecting the protein-ligand interaction into statistical or empirical "potentials" Such approaches however rely on parameter training using known protein-ligand binding structures or binding affinity measurements or both. Due to the limited size of the training set, the resulting scoring functions can be to too specialized (over-trained), thus have low transferability to targets and ligands that are structurally distinct from those in the training set 29 .

To eliminate these limitations, we report a novel scoring function MedusaScore for evaluating protein-ligand binding. MedusaScore describes the protein-ligand binding using physical interaction model. MedusaScore includes an explicit hydrogen-bonding model ³⁰ and EEF1 pairwise implicit solvent model³¹, which allows accurate modeling of the hydrogen-bonding and desolvation effect without large-scale MD simulations. Additionally, unlike other statistical and empirical scoring functions, MedusaScore does not depend on any specific parameter training on protein-ligand datasets, thereby maintaining the transferability over a wide range of potential drug candidates in the chemical space.

We test the performance of MedusaScore for docking decoy recognition and binding affinity prediction. Using the docking decoys datasets generated by Wang et al. 14 , we find that the recognition success rate for docking decoys is 82%, higher than that of 11 other scoring functions that are currently widely used in virtual screening, including LigScore, PLP⁷, PMF²⁴, LUDI²¹, F-Score⁸, G-Score⁹, D-Score¹¹, ChemScore²², Autodock¹⁰, DrugScore²⁵, and X-Score²⁶. The success rate can be further improved to 85% by consensus scoring with DrugScore²⁵. Using the PDBBind 2005 dataset 32 for the binding affinity prediction, we find that the MedusaScore showed a correlation coefficient of 0.60 and 0.61 for the core set and refined set (See Methods), respectively. This correlation is higher than what has been reported for 14 other scoring function using the same database 15 . Statistical analysis suggests that the entropic contribution may be the key component for further improvement of the accuracy of the binding affinity prediction.

Methods

Medusa force field

The Medusa force field 33 is a weighted sum of six energy terms:

 $E=W_{vdw_attr}E_{vdw_attr}+W_{vdw_rep}E_{vdw_rep}+W_{solv}E_{solv}+W_{bb_hbond}E_{bb_hbond}+W_{sc_hbond}E_{sc_hbond}+W_{bb_sc_hbond}E_{bb_sc_hbond}$

where E_{vdw_attr} , E_{vdw_rep} are the attractive and repulsive part of the van der Waals (VDW) interaction; E_{solv} is the solvation energy; E_{bb_hbond} , E_{sc_hbond} and $E_{bb_sc_hbond}$ are the hydrogen bond energies formed between backbone atoms, between side chains, and between backbone and side chains, respectively. The design of the force field is similar to that of the Rosetta force field 34 , which has also been widely used in protein folding and design. The VDW interaction model and parameters are adapted from CHARMM19 35 . The solvation model is the EEF1 implicit solvent model proposed by Lazaridis and Karplus 31 . We use the hydrogen bonding model proposed by Kortemme and Baker 36 . When evaluating the non-bonded interactions, we use a cutoff distance of 9.0 Å. The van der Waals repulsion (VDWR) potentials are implemented with linear extrapolation to dampen the fast increase of the potential as:

$$E_{vdw_rep} = \begin{cases} \sum\limits_{i,j>1} 4\varepsilon_{ij} \left[\left(\sigma_{ij}/r_{ij} \right)^{12} - \left(\sigma_{ij}/r_{ij} \right)^{6} \right], \alpha_{cutoff} \sigma_{ij} < r_{ij} \leq \sigma_{ij} \\ K_{slope} r_{ij} + 4\varepsilon_{ij} \left(\alpha_{cutoff}^{-12} - \alpha_{cutoff}^{-6} \right) - \alpha_{cutoff} K_{slope} \sigma_{ij}, r_{ij} \leq \alpha_{cutoff} \sigma_{ij} \\ Here, \alpha_{cutoff} = 0.92; K_{slope} = -24\varepsilon_{ij} \left(2\alpha_{cutoff}^{-13} - \alpha_{cutoff}^{-7} \right) / \sigma_{ij} \\ \varepsilon_{ij} = \sqrt{\varepsilon_{i}\varepsilon_{j}}; \sigma_{ij} = \sigma_{i} + \sigma_{j} \end{cases}$$

Here, r_{ij} is the distance between two atoms i and j. The energy parameters ε , σ are taken from the CHARMM19 force field of united atoms³⁵. Since the energy terms originate from different sources, a set of weighting parameters is assigned in order to balance their respective contributions. These weighting coefficients are trained to recapitulate the native amino acid sequence of 38 high-resolution crystal structures³³. The force field and the coefficients have been tested in various studies including experimental validations^{33,37-40}

New atom types and parameterizations

The original Medusa force field was designed to model atoms types occurring in proteins. Thus, to model small molecules for virtual screening, we extend the number of atom types as follows:

- **1.** For chemical groups that already exist in proteins, we keep the same atom types and parameters.
- 2. For new chemical groups, we define atom types based on (i) element types and hybridization, (ii) nearest-neighbor heavy atoms types, and (iii) second-nearest neighbor heavy atoms if they are charged. For the charged groups, we only consider carboxyl, phosphate and sulfone groups in current implementation.

Overall, we define 23 new atoms type in addition to the 38 existing atom types (Supplementary Table S1).

The VDW parameters are assigned according to atom sizes. For oxygen atoms, the VDW parameters are assigned the same as that of OC atom type in EEF1 ³¹, if they are charged, and same as OH, if they are not charged. The VDW parameters for nitrogen and sulfur atom types are the same as those used in CHARM19³⁵. The VDW parameters for P, F, Cl, Br and I are taken for Tripos TAFF force field. There are no new types for carbon atoms.

The extension of hydrogen bonding parameters is not needed since the parameterization of the model only depends on the hybridization types, on the explicit bond coordinates and on whether the hydrogen bonding is related to backbone³⁶. When applying the model, we treat all ligand atoms as protein side chain atoms, since usually there are no secondary structure constraints for ligand molecules.

There are two key parameters to extend in the EEF1 model: (1) the total solvation energy of the fully solvated atom ΔG_{free} and (2) the solvent volume V the atoms excludes 31 . The volume

parameters are assigned by considering the full volume of the atom and subtracting the overlap with neighboring bonded heavy atoms. The original ΔG_{free} parameters are taken from experimental solvation free energy measurements 31 . Since such measurements are not available for most small molecules, we assign these values according to their polarities. Following the original EEF1 model, we assign a large ΔG_{free} value of -20 kcal/mol for all charged groups. For other polar atoms, we use a linear relationship between the known partial charge and ΔG_{free} to assign the new ΔG_{free} values.

We use the same weights W_X as the original Medusa force field. Therefore, there is no additional training involved in force field parameters for developing the MedusaScore. The force field parameters are listed in Supplementary Table S1.

Scoring protocol

The scores are obtained from calculating the binding energy between the protein and the ligand using the extended Medusa force field. The protein coordinates are provided in a Protein DataBank (PDB) format and the small molecule coordinates in SYBYL mol2 format. Hydrogen atoms are required in the mol2 file to enable hybridization type assignment. All nonpolar hydrogen atoms are ignored in the protein input file since we use united-atom model. All polar hydrogen atoms are first reconstructed based on geometric bond constraints at physiological pH. Subsequently, the alternative positions of the rotatable polar hydrogen atoms are searched to optimize the protein-ligand hydrogen binding. To save computational time, the optimization is performed only for residues that are within 4 Å of the binding interface. All the hydrogen atoms in ligands are kept fixed during the optimization.

The computer program for MedusaScore is written in C++ and the simulation is performed on an Intel P4 2.4G Hz workstation running Gentoo Linux. The typical CPU time needed for the evaluation of a single protein-ligand complex is 0.11 s, of which 0.10 s is spent on parsing the input files.

Datasets

To benchmark our force field and scoring protocol, we use publicly available datasets that have been previously used to benchmark other scoring functions. This choice of a third party dataset allows comparison of MedusaScore with other scoring functions in an unbiased way. For discerning docking decoys, we use the dataset generated by Wang et al. ¹⁴, which has been used to compare docking accuracy for 11 scoring functions, including LigScore, PLP, PMF, LUDI, F-Score, G-Score, D-Score, ChemScore, Autodock, DrugScore and X-Score. For scoring, we use the protein pocket model in PDB format and the ligand coordinates in mol2 format.

For binding affinity prediction, we use the PDBBind database ³² (version 2005), which contains 1296 high quality complex structures (the refined set) and a subset of 288 non-redundant complexes (the core set). The latter contains structurally unrelated targets and ligands, and thus is more challenging for scoring function testing. Since our current force field does not take metal atoms into account, we eliminate all complexes that contain metal atoms within 4 Å of the ligand molecules (219 complexes total, Zn atoms in the carbonic anhydrase represent about one half of all cases). We also exclude complexes where a third molecule (mostly phosphate and sulfate) is bound to the same pocket, which corresponds to 16 complexes in the dataset. Finally, there are 4 complexes (PDB ID: 1duv, 1nw5, 1r6n, 2adm) containing atom types not modeled in the current force field implementation, which are also eliminated. After this filtering step, our dataset for benchmarking binding affinity prediction consists of 1057 complexes in the refined set and 243 complexes in the core set.

Results and Discussion

Docking decoy recognition

We apply the MedusaScore for the docking decoys of 100 protein-ligand complexes compiled by Wang et al 14 . Following the same criterion used by the authors 14 , we define a successful recognition when the best scoring (lowest binding energy) ligand decoy has a root-mean-square deviation (RMSD) less than 2 Å from the crystal structure (the RMSD is calculated only for ligand coordinates). We find that MedusaScore successfully recognizes the native-like poses for 82 of the entire 100 complexes, i.e., the success rate of MedusaScore is 82 %. This success rate is higher than that for eleven other scoring functions studied by Wang et al. 14 (see Table 1). The success rates of these scoring functions vary from 26 % (DSCORE) to 76 % (PLP). Interestingly, the force field-based scoring functions generally feature lower success rates than the other types of functions (see Table 1). The inclusion of EEF1 solvation model and the explicit hydrogen bond model is likely to contribute to the improved accuracy of the MedusaScore. An example of using MedusaScore in docking decoy recognition is shown in Figure 1 for a thermolysin inhibitor (PDB ID: 1tlp). From the 100 docking decoys generate using Autodock, MedusaScore correctly recognizes the native-pose while all other 11 scoring functions mistakenly picked non-native-like poses (RMSD > 6Å)

We further test the consensus scoring using MedusaScore and the other 11 scoring functions. We use a "rank-by-rank" strategy 41 , where the ranking of the decoys using both MedusaScore and other scoring functions are calculated and the decoy with the highest average rank is selected as the best scoring decoy. We find that consensus score with MedusaScore improves the decoy recognition rate for almost all other scoring functions except LIGSCORE, where the consensus scoring with MedusaScore has the same success rate as that of LIGSCORE alone.

On the other hand, we find that in most cases, consensus scoring decreases MedusaScore performance. The only exception is when MedusaScore is combined with PLP or DrugScore where the success rates are 83 % and 85 %, respectively. These success rates are higher than using MedusaScore alone (82 %). We attribute this improvement to the fact that PLP and DrugScore have significantly different energy potentials from ours. For example, DrugScore's energy function comprises of distance-dependent pairwise statistical potential and solvent-accessible surface dependent potential. None of these two potentials overlaps with the energy terms of MedusaScore. Due to these differences in the force field construction, some energy contributions that are ignored in MedusaScore may be realized in PLP or DurgeScore, thereby improving the accuracy. Another important fact is that both PLP and DurgScore also feature relatively high success rates. The MedusaScore is less likely to improve by consensus scoring with scoring functions that exhibit low recognition accuracies themselves. We also find that consensus scorings of MedusaScore with other force field-based scoring functions in general have lower success rates (Table 1). This observation likewise can be attributed to lacking of complementarity between MedusaScore and other force field-based scoring functions.

Consensus using 3 scoring functions does not further improve the success rate. We find the highest consensus scorings have success rate of 85 %, which are achieved from consensus score using MedusaScore, DurgScore and one of the third scoring functions (FSCORE, LUDI, or HMSScore, which is a scoring protocol from X-Score). Therefore, the consensus accuracy seems to be ultimately limited by the inherent inaccuracy of the individual scoring functions and no extra improvement can be obtained by further combination of the scoring functions.

Binding Affinity Prediction

Using the PDBBind database, we test the binding affinity prediction accuracy of MedusaScore. The correlation coefficient between the MedusaScore and experimental dissociation constant

 (pK_d) is 0.55 for the refined set and 0.56 for the core set. If we exclude the repulsive part of the van der Waals energies (VDWR) from the total score, the correlation improves to 0.61 and 0.60 for the refined set and core set, respectively (Figure 2). Similar observations have been reported in earlier studies 12 . This improved correlation by excluding VDWR may be due to clashes in some of the complex structures in the PDBBind 2005 datase. Besides subtracting the van der Waals repulsion directly, energy minimization of the complex structures should have the same effect. For simplicity, in the following analysis we only use the VDWR excluded MedusaScore.

Benchmarking studies have been reported ¹⁵ on an earlier version of PDBBind database (version 2002) for 14 scoring function including X-Score ²⁶, DrugScore ²⁵, D-Score ¹¹, PMF-Score ²⁴, G-Score ⁹, ChemScore ²², F-Score ⁸, LigScore, PLP ⁷, PMF ²⁴, LUDI ²¹, GoldScore ⁹, and HINT ⁴². The highest correlation coefficient is 0.566 using X-Score, lower than what we obtained using MedusaScore, albeit with a newer PDBBind database (version 2005) ³². To make a more objective comparison, we also calculate the MedusaScore using the 2002 version of the PDBBind Database. Similarly, we also exclude the complexes with metal or other heterogeneous atoms near the binding pocket. This procedure eliminates 181 out of the 800 complexes in the refined set. Using the VDWR-excluded scoring protocol, we find a correlation of 0.63 between the MedusaScore and the experimental binding affinity. This prediction accuracy is also significantly higher than that of the other 14 scoring functions that have been tested on the same dataset.

Although we do not use any binding affinity data for parameter training, MedusaScore still predicts the experimental values with reasonable accuracy. The robust performance over the various datasets suggests that MedusaScore likely grasps the crucial energetic component of protein-ligand binding. Since there is no training involved, MedusaScore should be applicable to a wide range of targets and ligands beyond those that have been tested in this study.

Size dependence

We further examine if the performance of MedusaScore depends on the ligand size. It has been reported that scoring functions tend to predict higher binding affinity for larger ligands, due to the inherent "stickiness" of the molecules 43 . To avoid any bias in the data analysis, we use the core set (See Methods) because it contains diverse structures. We divide the core set into three subsets, based on total number of heavy atoms (n) of the ligand molecule. The small, medium and large ligand size subsets correspond to $6 \le n \le 19$, $20 \le n \le 29$ and $30 \le n \le 70$ respectively, and contain 83, 82 and 78 complexes, respectively. We find that for the subset of small ligands, our scoring function features the best correlation with experimental data — the Pearson correlation coefficient is 0.63, higher than the average correlation over the whole dataset (Figure 3). The correlation decreases for larger ligands — the correlation coefficients for the medium and large ligands are 0.52 and 0.37, respectively, lower than the average over the whole dataset. In general, our observations agree with the previously described tendency of larger molecules to have lower scores 43 .

When we divide the dataset according to the number of heavy atoms of the ligand that are in contact with the protein (n_c) , we find a better correlation between MedusaScore estimates and the experimental data. The three subsets, defined by $1 \le n_c \le 6$, $7 \le n_c \le 9$ and $10 \le n_c \le 25$, contain 88, 72 and 83 protein-ligand complexes, respectively. The contact is defined when the ligand atom is within 3.5 Å of any protein heavy atom. We find that our scoring function is most accurate for the subset with the least contact atoms $(1 \le n_c \le 6)$, for which the correlation coefficient is 0.74. The correlation decreases to 0.57 and 0.42 for the subsets of $7 \le n_c \le 9$ and $10 \le n_c \le 25$, respectively. Similarly to the previous case, we observe systematic overestimation of the binding affinity for ligands with larger n_c .

Clearly MedusaScore can be utilized more confidently for smaller-sized ligands or ligands with a smaller number of contacts with proteins. This is partly due to the inaccuracies in the force field, which tend to accumulate with the increase of the ligand size or the number of contacts with the proteins. However, these force field inaccuracies alone cannot explain the systematic overestimation of binding affinities for ligands with larger n or n_c . The experimental binding affinities have less significant dependence on n or n_c as shown in Figures 3 and 4, because the favorable binding enthalpies for larger molecules are often compensated by the larger entropic loss upon binding 44 . This enthalpy-entropy compensation effects are not considered in MedusaScore, which explains the over-estimation of binding affinity.

It has been suggested that entropic contribution may play an important role in ligand binding and that entropic contribution can be as large as the enthalpy 45 , and therefore may need to be properly evaluated to achieve a better accuracy in protein-ligand binding scoring. Furthermore, close contacts introduce significant fluctuations in the total energy due to the sensitivity/volatility of the VDWR energy term. These factors might be important for further MedusaScore improvement.

VDWR Terms and Energy Minimization

We have shown that excluding the VDWR term from MedusaScore in general results in slightly improved binding affinity prediction. We further test whether this observation is due to clashes in the molecular complexes and whether energy minimization can improve the prediction accuracy. We examine the VDWR energies in the core set and find that there are five complexes having exceptionally large VDWR energies (> 25 kcal/mol), compared with the rest having an average VDWR energy of 4.7 kcal/mol and a standard deviation of 8.9 kcal/mol. After excluding these outliers from the core set, the correlation coefficient between the MedusaScore prediction (with all energy terms) and experimental binding affinity is 0.59. Further removal of the VDWR term only marginally improves the correlation to 0.60. We attempt to minimize the structures by allowing local rigid body movement of the ligand and side chain movement of the protein. Such relaxation reduces VDWR energies in general (by $\sim 3-6 \text{ kcal/mol}$, see Supplementary Table S2), but fails to reduce the high VDWR energies for the outliners effectively. As a result, the overall correlation is not improved after the minimization.

Therefore, our results demonstrate that without performing explicit minimization, removing the VDWR term, in fact, makes the scoring function more robust to structure imperfections. This strategy might be necessary for virtual screening because it saves costly computational time for structure minimizations. Unphysically close atom contacts in the complex structures can be detected by applying a quick atomic distance filter before using the scoring function.

Comparison with RosettaLigand

It is interesting to compare the performance of MedusaScore with the scoring function of RosettaLigand 12 since the underling force fields behind the two scoring functions have similar energy terms. To our best knowledge, the only published benchmark results for RosettaLigand are in the original publication 12 , where the RosettaLigand's scoring function was used for binding affinity prediction on LPDB database 46 . The authors found a correlation of 0.63 between the experimental and predicted binding affinities.

In order to compare the performance of MedusaScore with RosettaLigand, we test the performance of MedusaScore using the LPDB database. The current LPDB database contains 262 complexes. Following the same protocol as described in Methods, we eliminate complexes with unsupported atom types or having metal and other additional hetero molecules near the binding interface. After the preprocessing, we obtain 196 complexes for benchmarking. Applying MedusaScore on this dataset results in correlations of 0.63 and 0.64 with and without

the VDWR term, respectively. These correlations are comparable with the previously published results from RosettaLigand 12 .

Although RosettaLigand and MedusaScore features similar accuracy in terms of binding affinity prediction, there are several notable differences between the two scoring functions: (1) RosettaLigand uses Tripos mol2 based atom types for ligands, while MedusaScore uses extension of CHARMM19 atom types. (2) The weighing coefficients have been retrained in RosettaLigands using 100 protein-ligand structures ¹², while we do not retrain any weighing coefficient in MedusaScore.

Conclusion

We have developed a scoring function for protein-ligand interaction by extending the Medusa force field and design suite. Benchmarking using available datasets show superior performance of the MedusaScore for both docking decoy recognition and binding affinity prediction. Since the MedusaScore does not rely on parameter training using protein-ligand binding data, it is transferable to targets and small molecules beyond the tested datasets. Therefore, we expect the MedusaScore to have wide application in virtual screening of novel small molecule drug candidates.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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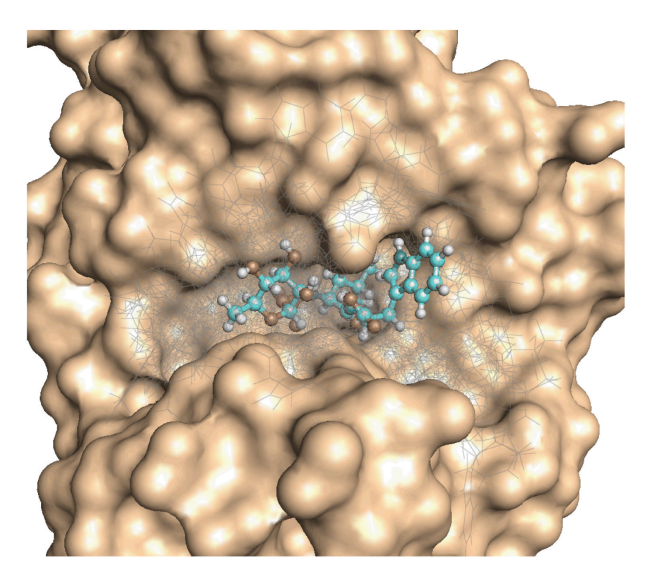


Figure 1. An example of docking decoy recognition using MedusaScore The native docking pose (ball-and-stick) for a inhibitor against thermolysin protein (PDB ID: 1tlp) is correctly recognized using MedusaScore from the 100 docking decoys (gray lines) generate by Wang et al. 14 using Autodock 10 .

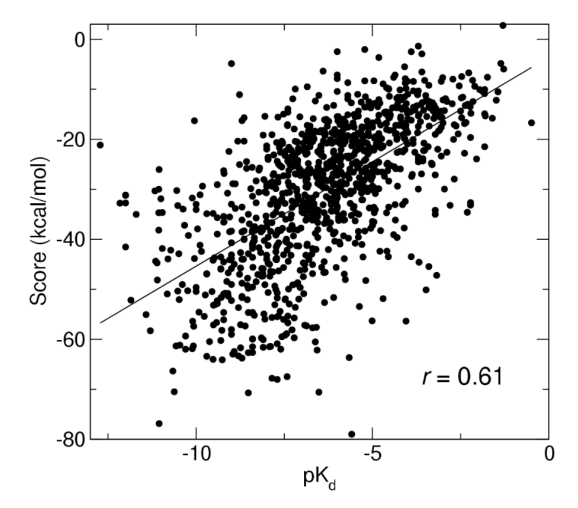


Figure 2. Scatter plot of the MedusaScore predictions (without VDWR) vs. the experimental dissociation constant pK_d for the refined set The Pearson correlation coefficient is 0.61. The solid line corresponds to a linear regression

fit (y = 4.18x - 3.59).

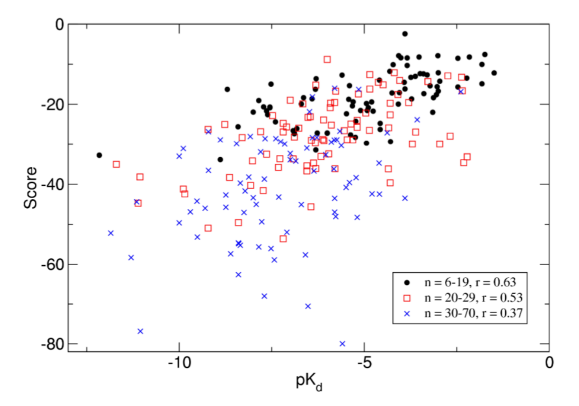


Figure 3. Scatter plot of the MedusaScore vs. the experimental dissociation constant pK_d for the core set, categorized based on the number of heavy atoms in the ligand (n) The prediction accuracy is higher (r=0.63) for small ligands, than for medium and large sized ligands.

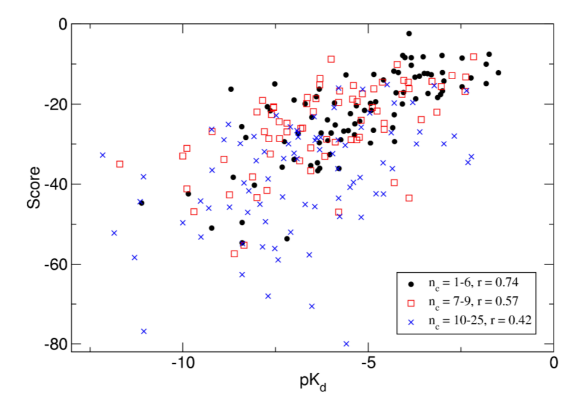


Figure 4. Scatter plot of the MedusaScore vs. the experimental dissociation constant pK_d for the core set, categorized based on the number of ligand heavy atoms in contact with protein (n_c) We define a contact when a ligand heavy atom is within 3.5 Å of any protein heavy atom. The prediction accuracy is higher (r = 0.74) for ligands with fewer contacts with the proteins than for those with more extensive contact.

Table 1 Success rates of MedusaScore and other 11 scoring functions for docking decoy recognition

The scoring functions are tested on docking decoy dataset consisting of $100 \text{ complexes}^{14}$. The success rate is defined when the best scoring decoy ligand has RMSD less than 2 Å from the crystal structure. The MedusaScore has the highest success rate of all the tested functions (82 %). Consensus scoring with the MedusaScore improves other scoring functions. The combination of the MedusaScore with PLP or with the DrugScore gives a success rate of 83 % and 85 %, respectively (highlighted by the bold font), higher than the MedusaScore alone.

Scoring function		Success rate	
Туре	Name	Single Scoring	Consensus with Medusa
Force field	MedusaScore	82	-
	Autodock	61	71
	GScore	42	74
	DScore	26	72
	PLP	76	83
	FScore	74	79
Empirical	LigScore	74	74
	LUDI	67	82
	X-Score	65	78
	ChemScore	35	60
Statistical potential	DrugScore	71	85
	PMF	52	70