## Learning from Docked Ligands: Ligand-Based Features Rescue Structure-Based Scoring Functions When Trained On Docked Poses

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#### Abstract

Machine learning scoring functions for protein-ligand binding affinity have been found to consistently outperform classical scoring functions when trained and tested on crystal structures of bound protein-ligand complexes. However, it is less clear how these methods perform when applied to docked poses of complexes.

We explore how the use of docked, rather than crystallographic, poses for both training and testing affects the performance of machine learning scoring functions. Using the PDBbind Core Sets as benchmarks, we show that the performance of a structure-based machine learning scoring function trained and tested on docked poses is lower than that of the same scoring function trained and tested on crystallographic poses. We construct a hybrid scoring function by combining both structure-based and ligand-based features, and show that its ability to predict binding affinity using docked poses is comparable to that of purely structure-based scoring functions trained and tested on crystal poses. Despite strong performance on docked poses of the PDBbind Core Sets, we find that our hybrid scoring function fails to generalise to anew data set, demonstrating the need for improved scoring functions and additional validation benchmarks. Code and data to reproduce our results are available from https://github.com/oxpig/learning-from-docked-poses.

### Introduction

Rapidly prioritising which compounds to make is a key question in early-stage drug discoverv<sup>1</sup>. When the structure of the target is known, one commonly-used approach is highthroughput protein-ligand docking,<sup>2,3</sup> which uses scoring functions to rank compounds by their predicted affinity for the target protein. Such scoring functions must make several assumptions that trade biophysical accuracy for speed, and rely predominantly on structurebased features to quantify the various protein-ligand interactions. It is common practice to use X-ray crystallographic structures of bound protein-ligand complexes when training and testing these models to predict protein-ligand binding affinity. This is reasonable, as it isolates the task of binding affinity prediction from structural errors that might be introduced as a result of inaccurate or incorrect ligand pose prediction, or due to the rigid receptor assumption that is often used in docking.<sup>4–6</sup> However, in a real-world drug discovery scenario, it is highly unlikely that a crystal structure of the bound protein-ligand complex will be available for every ligand or class of ligands of interest. Instead, protein-ligand docking is often used to predict the binding mode of each ligand within the binding pocket of a protein target. Thus, for scoring functions to be relied upon in prospective screens, they will need to show good predictive performance not only on experimentally-determined binding poses, but also on docked poses.

Although less common than the use of crystal structures, there are a few reported studies of the effect of using docked poses in training and testing scoring functions. Durrant and McCammon<sup>7</sup> used both crystallographic and docked poses in the development and evaluation of their scoring function, NNScore 2.0, and noted that the optimal choice of docking protocol was highly system-dependent, but did not examine the difference in performance between

training using only crystallographic or only docked poses of the same complexes. Zilian and Sotriffer<sup>8</sup> trained SFCScore<sub>RF</sub> using crystal poses from the PDBbind database,  $^{9,10}$  and validated on a combination of crystal poses from the PDBbind database and docked poses from the CSAR–NRC HiQ<sup>11</sup> and the CSAR 2012 benchmarks.<sup>12</sup> SFCScore<sub>RF</sub> demonstrated strong performance on crystallographic poses of the PDBbind 2007 Core Set, with a Pearson correlation coefficient between the predicted and experimental binding affinity of 0.779; and on docked poses of the CSAR–NRC HiQ test set, with a Pearson correlation coefficient between predicted and experimental binding affinity of 0.730. Performance on the CSAR 2012 benchmark was highly target-dependent, with the authors attributing poor performance on the kinase targets CHK1 and ERK2 to the poor quality of docked poses. More recently, Jimenez et al.<sup>13</sup> validated  $K_{DEEP}$  using both crystal poses and docked poses but, as in the earlier studies, used docked poses only when crystal structures were unavailable, so the impact of the use of docked poses in place of crystal structures was not determined. Li et al.<sup>14</sup> investigated how the use of docked poses in place of crystal poses affected the performance of the AutoDock Vina scoring function, RF-Score, and RF-Score v3. They reported that, contrary to what might be expected, using docked poses in place of crystal poses had only a small effect on the accuracy of their binding affinity predictions; for example, the Spearman rank correlation coefficient between predicted and experimental binding affinity attained by RF-Score v3 on the PDBbind 2013 Core Set dropped from 0.662 (crystal poses) to 0.633 (docked poses). The authors also reported that this drop in performance was reduced by training RF-Score v3 on docked poses instead of crystal poses, resulting in a Spearman rank correlation coefficient between predicted and experimental binding affinity of 0.643 on the PDBbind 2013 Core Set, suggesting that training on docked poses can make a scoring function less susceptible to errors when subsequently tested on docked poses.

Here, we describe a new approach to compensate for potential errors introduced into protein-ligand binding affinity prediction through the use of docked (instead of crystallographic) poses. Previously,<sup>15</sup> we reported that the inclusion of pose-independent ligand-

based features improved the performance of several machine-learning scoring functions in predicting protein-ligand binding affinity when trained on crystallographic binding modes of the PDBbind database,<sup>9</sup> and tested on the Comparative Assessment of Scoring Functions (CASF) benchmarks.<sup>4-6</sup> We also found that ligand-based features are predictive of the average binding affinity of a ligand for its protein targets in PDBbind<sup>15</sup>. Motivated by this, and the fact that ligand-based features are not affected by errors introduced by the use of docked poses, we investigated how the inclusion of a rich set of ligand-based features as inputs to a machine learning scoring function affects its performance when trained and tested on docked poses. Firstly, we investigated the effect of training and testing scoring functions using only docked poses generated by Smina,<sup>16</sup> a fork of AutoDock Vina,<sup>17</sup> as opposed to X-ray crystallographic poses, on the resulting scoring functions. We found that pose prediction errors are common when re-docking the ligands in the PDB bind database into their corresponding protein co-crystal structures, and that even when accurate poses were generated, the Smina scoring function often failed to rank these native-like poses higher than non-native ones. We also found that training and testing machine learning scoring functions on docked poses degrades scoring function performance relative to those trained and tested on crystallographic poses even when there is a high level of similarity between proteins in the training and test sets. Augmenting the input features with rapidly computed ligand-based molecular descriptors results in a greater gain in performance when using docked poses, than when using crystal poses for training and testing.

Finally, we construct a new data set consisting of six of the eight protein targets from the DUD-E Diverse Subset,<sup>18</sup> for which ligand binding affinity data were available in the ChEMBL database, version 25,<sup>19</sup>. We find that our models perform well when trained and tested on docked poses of ligands for a single target; but poorly when trained and tested on docked poses for different targets. While there is clearly some way to go in order to create truly generalised machine learning scoring functions, our results suggest that the inclusion of additional ligand-based input features in a scoring function helps to correct for potential errors introduced by docking. This thus greatly expands the set of targets and ligands that can be used to train new scoring functions, and will help to generate better predictions of protein-ligand binding affinities.

## Methods

#### Data

#### **PDBbind Training Set**

We used the PDBbind 2018 Refined Set<sup>10</sup> of high-quality protein-ligand structures and binding affinities as our source of training data. This consists of 4,463 crystal structures of protein-ligand complexes from the PDB<sup>20</sup> with experimentally-determined binding affinity values. For our training set we selected all the structures in the PDBbind 2018 Refined Set, but excluded 481 structures for which either the docking failed or features could not be computed; we also excluded any structures that were also present in the PDBbind Test Set (described below). This resulted in a training set of 3,752 high quality protein-ligand complexes with corresponding binding affinity data, which we refer to as the "PDBbind Training Set".

We used the PDBbind 2018 Refined Set rather than the larger PDBbind 2018 General Set<sup>10</sup> for two reasons. Firstly, structures in the Refined Set are subject to strict quality controls, including a resolution of 2.5Å or better and no missing side chains, reducing the risk of errors in the docking results that might be introduced by the use of inaccurate crystal structures. Secondly, while several authors have reported improved performance on the PDBbind Core Sets when training on the larger General Set instead of the Refined Set,<sup>21,22</sup> we observed previously<sup>15</sup> that this can be attributed to increased representation of the Core Set proteins in the training set: when proteins with high sequence identity to those in the test set were excluded from the training set, there was little difference in performance between

training on the General Set and training on the Refined Set.<sup>15</sup>

#### PDBbind Test Set

To evaluate the performance of our scoring functions, we constructed a test set by combining the structures in the PDBbind 2007, 2013, and 2016 Core Sets, which correspond to the test sets used in the Comparative Assessment of Scoring Functions (CASF) 2009, 2013, and 2016, respectively.<sup>4–6</sup> Combining these three PDBbind Core Sets, and removing 30 structures for which features could not be computed due to RDKit version 2019.09.1<sup>23</sup> failing to sanitise the ligand, resulted in a test set of 525 protein-ligand complexes, which we refer to as the "PDBbind Test Set". Previously, we found that using this combined test set of unique protein-ligand complexes in place of the smaller Core Sets resulted in a better (i.e. narrower) confidence interval for the performance metric.<sup>15</sup>

#### Updated DUD-E Diverse Subset

We constructed an Updated DUD-E<sup>18</sup> Diverse Subset for six of the eight diverse protein targets for which  $K_i$  binding affinity data could be found in version 25 of the ChEMBL database.<sup>19</sup>. These targets were: (1) serine/threonine-protein kinase AKT (AKT1); (2) cytochrome P450 3A4 (CP3A4); (3) glucocorticoid receptor (GCR); (4) HIV-1 protease (HIVPR); (5) HIV-1 reverse transcriptase (HIVRT); and (6) kinesin-like protein 1 (KIF11). Two of the eight targets in the DUD-E Diverse Set,  $\beta$ -lactamase (AMPC) and C-X-C chemokine receptor type 4 (CXCR4), which were the smallest sets in the original DUD-E Diverse Subset, did not have any ligands with recorded  $K_i$  measurements in ChEMBL at the time of writing, and so were excluded.

For each target, we queried ChEMBL version 25 for ligands that bind to that target and which had one or more measurement of  $K_i$ . We used only  $K_i$  data and excluded measurements such as IC<sub>50</sub> so as not to conflate different types of data. We trained our models using the corresponding pChEMBL value, which is the negative base-10 logarithm of the binding constant reported by ChEMBL, equivalent to the pK value.<sup>24</sup> Duplicate pChEMBL values for a protein-ligand pair were removed; for ligands with different pChEMBL values for the same target, we used the arithmetic mean of the pChEMBL value. The DUD-E abbreviations for the names of the six targets for which  $K_i$  data were available in ChEMBL, together with the PDB code of the structure provided by DUD-E and the number of ligands for each target, are shown in Supporting Information Table 1.

The majority of the data for HIVRT and CP3A4 span only four orders of magnitude, with pChEMBL values ranging from 4 to 8. The data for GCR cover a slightly larger range, with pChEMBL values ranging from 4.7 to 10, while the data for AKT1, HIVPT, and KIF11 span at least six orders of magnitude. The binding affinities all lie within the range of values represented in the PDBbind 2018 Refined Set, so a Random Forest (RF) model trained on PDB bind data could be expected to interpolate successfully. The distributions of the pChEMBL values for the six targets are shown in Supporting Information Figure 1. We performed three different validation experiments using the Updated DUD-E Diverse Subset. First, to explore the effect of using docked poses in place of crystallographic poses, we trained on docked poses (see "Docking Protocol") of the PDBbind Training Set and tested on docked poses of all of the ligands obtained from ChEMBL for each target. Second, we investigated whether our models could learn from docked poses obtained by docking ligands taken from ChEMBL into a single structure of a protein target, rather than using the co-crystal cognate structure of the protein each ligand. To do this, for each target we randomly selected 80%of the ligands obtained from ChEMBL for that target to use as a training set. The resulting model was then tested on the remaining 20% of the ligands for that target (so-called "Intratarget training"). Third, we investigated whether our models could generalise to ligands for a previously-unseen target when trained upon data other than the PDBbind Refined Set. For each target we trained on all ligands known to bind to the other five targets, and tested on the held-out target's set of ligands (so-called "Inter-target training").

#### **Docking Protocol**

All docking calculations were performed using Smina<sup>16</sup> (version November 9 2017), a fork of AutoDock Vina.<sup>17</sup> Protein and ligand structures were prepared for docking using the following protocol. For each ligand, an initial conformer was generated using the ETKDG method<sup>25</sup> implemented in RDKit version 2019.09.1.<sup>23</sup> For the PDBbind data, this ensures the docking could not be biased by starting with the crystallographic bond lengths, bond angles, and torsions of the ligand. PDBQT files for both the receptor and ligand were generated using OpenBabel.<sup>26</sup> We used the default parameters of Smina, with the following exceptions:  $autobox_add = 8$ ; exhaustiveness = 20; and  $num_modes = 20$ . For each protein-ligand complex from PDBbind, the centre of search space was defined by passing the crystallographic binding mode of the ligand using the *autobox\_ligand* parameter. For each ligand, up to 20 diverse docked poses were generated by Smina. In addition to docking, we also performed a local AutoDock Vina energy minimisation of the crystallographic binding mode of the ligand provided by PDB bind using Smina to generate a single near-native docked pose. To perform the minimisation, we used Smina's *minimize* option with default parameters. For the Updated DUD-E Diverse Subset, we docked the ligands into the PDB structure provided by DUD-E for each target, using the crystallographic ligand binding pose provided by DUD-E to define the centre of the search space.

The quality of a docked pose was assessed by computing the root-mean-squared deviation (RMSD) of the coordinates of the atoms of the ligand's docked pose with respect to the coordinates of the ligand's atoms in the crystallographic pose. To ensure we correctly accounted for symmetry when computing the RMSD between two conformers, we identified symmetrically equivalent permutations of the atomic indices of a molecule by performing a substructure match of the molecule against itself using RDKit version 2019.09.1. We then applied these permutations to the indices of the atoms and re-computed the RMSD of the docked and crystallographic structures for each permutation. The lowest computed RMSD value was then taken as the RMSD of that docked pose.

#### Scoring Function Construction

We used the Random Forest (RF)<sup>27</sup> as implemented in Scikit-Learn version 0.22.0<sup>28</sup> as our learning algorithm, as our previous results demonstrated that it consistently outperformed other tested machine learning methods.<sup>15</sup> We built three models to predict the binding affinity of a protein-ligand complex that differed in the types of input features used: a purely ligand-based (LB) model; a traditional structure-based (SB) model using proteinligand intermolecular features; and a hybrid (HB) model consisting of both ligand-based and structure-based features.

For the input features of the ligand-based (LB) model, we used a set of pose-independent molecular descriptors computed for each ligand using RDKit version 2019.09.1. These descriptors are conformation-independent and may be categorised as either (computed) experimental bulk properties (such as molar refractivity or logP) or theoretical descriptors derived from a symbolic representation of the molecule. The theoretical descriptors may be further categorised according to the dimensionality of the representation of the molecule from which they are derived. The conformer-independent descriptors we consider are either 1-D compositional properties (such as heavy atom counts, bonds counts, and molecular weight) or 2-D topological properties (such as fragment counts, topological polar surface area, and connectivity index). Any features with zero variance across the training data set, or that were null-valued (*i.e.* infinite or not computable) were excluded. We removed the Ipc index<sup>29</sup> as it produced extreme numerical values for larger molecules that were too large to be represented as 32-bit floats. This resulted in 185 ligand-based features, and the full list of features is given in the Supporting Information, under "RDKit Features".

Our structure-based (SB) model uses the features of the Random Forest-based scoring function RF-Score v3<sup>30</sup>. Six of these features are the same six terms used by the AutoDock Vina scoring function (five empirical force-field-like potentials derived from the interactions between protein and ligand atoms, and the number of rotatable bonds in the ligand). The remaining 36 features are the counts of pairwise interactions between protein and ligand atoms within 12Å of each other, for example, the number of protein carbon-ligand nitrogen pairs. Four elements (C, N, O, and S) are considered in the protein and nine elements (C, N, O, F, P, S, Cl, Br, and I) in the ligand. These features were calculated using the Open Drug Discovery Toolkit version 0.6.<sup>31</sup>

For the hybrid (HB) model, we used the features of both the ligand-based and structurebased models as inputs and again trained Random Forest models using the training set described above.

#### Model Training and Testing

#### **PDBbind Validation**

To investigate the effect of using docked or crystallographic ligand binding modes on binding affinity prediction, we performed a five-fold cross-validation on the PDBbind Training Set. Four approaches to training our scoring functions were compared, training on structure-based features derived from: (i) the crystallographic pose of the ligand; (ii) a single docked pose obtained by performing local minimisation of the ligand using Smina; (iii) a single docked pose, ranked highest by Smina; and (iv) multiple docked poses for each ligand, in this case up to 20 diverse poses per ligand generated by Smina. In case (iv), each pose was labelled with the same experimental binding affinity value during training.

We tested three strategies for predicting the binding affinity of a ligand using its docked poses: (i) predicting the binding affinity using the pose ranked highest by Smina; (ii) predicting the binding affinity for each docked pose and taking the highest; and (iii) predicting the binding affinity for each docked pose and taking the arithmetic mean.

#### Updated DUD-E Diverse Subset

Using the six targets of the Updated DUD-E Diverse Subset, we applied three different approaches to training and validation. First, models were trained on docked poses of the PDBbind Training Set and tested on docked poses of the Updated DUD-E Diverse Subset. Second, we performed an "inter-target" validation in which each of the six targets in turn was held out as a test set with the remaining five targets forming the training set. Third, we performed an "intra-target" validation in which we randomly selected 20% of the ligands for each target to be a validation set, and trained on the remaining 80% of the ligands for that target. We then repeated both the inter-target and intra-target validation experiments by combining the DUD-E/ChEMBL data with the PDBbind Training Set.

The performance of each scoring function was evaluated by computing the Pearson correlation coefficient between the predicted and experimentally-determined values of the proteinligand binding affinity, expressed as  $pK_i$  values.

## **Results and Discussion**

#### Accuracy of docked poses

We first evaluated the quality of the predicted binding poses generated by Smina for the 4,277 ligands in the PDBbind Training Set (3,752) and PDBbind Test Set (525). For each ligand, up to 20 diverse poses were generated. Only 1,357 ( $\approx 32\%$ ) of the ligands re-docked by Smina had at least one pose with with an RMSD less than 2Å with respect to the crystallographic coordinates. For 1,003 ligands ( $\approx 23\%$ ), every pose generated by Smina had an RMSD greater than 4Å. The distribution of the lowest RMSD docked pose for each complex is shown in Supporting Information Figure 2. The AutoDock Vina scoring function used by Smina often failed to rank a near-native ("good") pose first even when one was generated by Smina: of the 1,357 complexes with at least one pose with an RMSD greater than 2Å, about half (691) of the top-ranked poses returned for Smina had an RMSD greater than 2Å, and about a quarter (371) of these had an RMSD of greater than 4Å. This agrees with previous results that have shown that docking scoring functions are not always a reliable way of selecting the best pose from a set of putative binding poses.<sup>32</sup> One well documented source of error in docking is the difficulty of sampling the conformational space of large,

highly flexible molecules. However, we found little correlation between either the size or the flexibility of the ligand and the RMSD of the best docked pose (Supporting Information Figure 3).

# Effect of the quality of the docked poses on model training and testing

As there was considerable variation in the accuracy of the docked poses generated by Smina, we investigated how the quality of the docked pose, or poses, used to train and test our models affected binding affinity prediction. For this, we trained and tested each model using five-fold cross-validation of the PDBbind Training Set. In our first set of experiments, to determine the influence of the ligand's pose on model performance, we trained the models using a single minimised pose (obtained by minimising the crystallographic binding pose using Smina) of each ligand in the training folds and tested on all of the (up to) 20 docked poses of each ligand in the test fold. We performed three experiments investigating strategies for making a single prediction of binding affinity using multiple docked poses of the test ligand. These involved using our models to predict the binding affinity for: (1.1) the pose ranked first by Smina; (1.2) all (up to) 20 docked poses, taking the highest value as the predicted affinity; (1.3) all (up to) 20 poses and taking the mean score as the predicted affinity; and (1.4) training and testing using the crystallographic coordinates of each ligand, as a control. Minimised crystal poses were used during training for experiments (1.1)-(1.3) to control for possible errors due to poor docking while ensuring that, like the docked poses for the test set, the training poses were optimised with respect to the Smina scoring function.

Table 1 compares the average Pearson correlation coefficient achieved in each experiment under five-fold cross validation using the three docking-based scoring strategies (experiments 1.1-1.3) with training and testing on crystal poses (experiment 1.4). The performance of the ligand-based (LB) model is shown for comparison; its performance is necessarily the same regardless of scoring strategy, as the ligand-based features are pose-independent. Both the structure-based (SB) model and hybrid (HB) model (combination of LB and DB features) perform worse when trained and tested on docked ligand poses instead of the experimentallydetermined crystallographic poses, regardless of the scoring strategy. Using the mean predicted affinity across a set of docked poses yields particularly poor results when compared to training and testing on crystal poses, with the Pearson correlation coefficient between predicted and experimentally-determined binding affinity dropping from 0.746 to 0.604 for the SB model, and from 0.768 to 0.676 for the HB model. Testing on the pose ranked best by Smina results in a smaller drop in Pearson correlation coefficient than when taking the mean of the predicted affinities across multiple poses: to 0.659 for the SB model and to 0.714 for the HB model. The large drop in performance of the SB model is in contrast with the results of Li *et al.*, who found that training and testing on docked poses resulted in only a small difference in performance<sup>14</sup>. One possible explanation for this difference is that by training only on minimised poses the models only see docked poses that are very close to the crystal pose, and so may not be capable of extrapolating to the less accurate docked poses used for testing.

Table 1: Pearson correlation coefficient between predicted and experimental binding affinity under five-fold cross-validation on the PDBbind Training Set. Random Forest models were trained using ligand-based (LB) features, structure-based (SB) features derived from a single minimised pose, or using both LB and SB features (HB), and tested on up to 20 docked poses, or the crystallographic pose of each ligand. The Pearson correlation coefficient was computed between the experimental and predicted binding affinity values obtained in four experiments: (1.1) scoring the pose ranked best by Smina, (1.2) scoring all poses and taking the maximum score, (1.3) scoring all poses and taking the mean score, and (1.4) training and testing using the experimentally-determined binding pose. As expected, the LB model is insensitive to the docked ligand pose(s). The best performance for each scoring strategy is underlined.

Experiment	Train Pose(s)	Test Pose(s)	LB Model	SB Model	HB Model
1.1	Minimised	Тор	<u>0.719</u>	0.659	0.714
1.2	Minimised	All (max)	0.719	0.682	0.732
1.3	Minimised	All (mean)	<u>0.719</u>	0.604	0.676
1.4	Crystal	Crystal	0.719	0.746	0.768

Having found that for models that used structure-based features, testing on minimised crystallographic poses and testing on docked poses results in lower performance than when training and testing on crystal poses, we next investigated whether training on docked poses instead of minimised crystal poses would enable the models to better generalise to docked poses. We thus repeated experiments 1.1, 1.2, and 1.3, but this time trained on the pose ranked best by Smina, rather than the minimised crystallographic pose. The results are shown in Table 2. In all cases, our models perform better when trained on docked poses rather than minimised crystallographic poses. This difference is marginal when taking the highest predicted affinity of the docked poses of the test ligands (2.2), but noticeable when using just the docked pose ranked best by Smina (2.1), or when taking the mean predicted of the poses (2.3). This suggests that training on docked poses helps our models to generalise to docked poses of varying accuracy, and that when presented with multiple docked poses, the highest score assigned by our models is likely the most accurate, reflective of the actual protein-ligand binding affinity.

Regardless of the types of ligand poses used for training and testing, the LB model consistently outperformed the SB model under cross-validation. Even better performance was obtained using the HB model when trained on the docked pose ranked highest by Smina, and using this to predict the affinities across all poses and taking the best binding affinity; indeed, this performance (2.2;  $\rho_{\rm P} = 0.744$ ) was comparable to our SB model when trained and tested on crystal poses (1.4;  $\rho_{\rm P} = 0.746$ ), suggesting that augmenting structure-based features with ligand-based features in a machine learning-based scoring function can help to correct for errors that might be introduced by the use of docked poses. Table 2: Pearson correlation coefficient between predicted and experimental binding affinity under five-fold cross-validation on the PDBbind Training Set. Random Forest models were trained using the docked pose ranked highest by Smina and tested on up to 20 docked poses for each ligand. The Pearson correlation coefficient was computed using affinity predictions obtained in four ways: (2.1) scoring the pose ranked best by Smina (top pose); (2.2) scoring all poses and taking the maximum (all poses (max score)); (2.3) scoring all poses and taking the mean (all poses (mean score)); and (1.4) training and testing on the crystal pose of each ligand (crystal pose). As expected, the LB model is insensitive to the docked ligand pose(s). The best performance for each scoring strategy is underlined.

Experiment	Train Pose(s)	Test Pose(s)	LB Model	SB Model	HB Model
2.1	Тор	Тор	0.719	0.676	0.738
2.2	Тор	All (max)	0.719	0.687	0.744
2.3	Тор	All (mean)	0.719	0.643	0.725
1.4	Crystal	Crystal	0.719	0.746	<u>0.768</u>

#### Effect of training using multiple poses

As training on a single docked pose for each ligand was more effective than training on minimised crystal poses, we next investigated how training on multiple docked poses for each ligand affected model performance. To do this, we repeated the cross-validation experiment described above, this time in experiments 3.1, 3.2, and 3.3 training on all docked poses for each ligand, using the same binding affinity value for each pose of a ligand. We used our ML models to score all poses of the ligands in the test fold and took the maximum predicted binding affinity as the predicted value for that ligand. To control for the effect of substantially increasing the size of the training set, we repeated this cross-validation, but for each training ligand used the same number of copies of the pose ranked top by Smina as the number of diverse docked poses output for that ligand by Smina (up to 20). Table 3 shows the mean Pearson correlation coefficient achieved by each model over the five crossvalidation folds, when trained on multiple diverse or redundant poses. Overall, there is little difference between training on a single pose and training on multiple poses or redundant poses, with the SB model performing slightly better when trained on all poses instead of one pose ( $\rho_{\rm P} = 0.699$  vs  $\rho_{\rm P} = 0.676$ ), and the HB model performing slightly worse ( $\rho_{\rm P} = 0.723$  vs.  $\rho_{\rm P} = 0.738$ ). Similarly, training on multiple copies of the pose ranked highest by Smina does not significantly affect performance, indicating that the models are not adversely affected by redundancy in the training data. These results, together with those in Table 1 and Table 2, suggest that when using Random Forest for protein-ligand binding affinity prediction, it is important for the model to see examples of docked poses of varying quality, but using a variety of example poses for each complex is not necessary.

Table 3: Mean Pearson correlation coefficients between predicted and experimental binding affinity under five-fold cross-validation on the PDBbind Training Set. For each ligand, models were trained using either the pose ranked highest by Smina (top pose), all diverse poses, or redundant copies of the pose ranked highest by Smina. Predictions were made for the test fold by scoring all docked poses of each ligand and taking the highest score. The best performance for each experiment is underlined.

Experiment	Train Pose(s)	Test Pose(s)	LB Model	SB Model	HB Model
3.1	Тор	All (max)	0.719	0.676	0.738
3.2	Diverse	All (max)	0.713	0.699	<u>0.723</u>
3.3	Redundant	All (max)	0.713	0.689	0.745

#### Validation on PDBbind Test Set

We next validated our LB, SB, and HB models on the PDBbind Test Set. Each model was trained on the PDBbind Training Set. We used the docked pose ranked highest by Smina for each ligand in the training set and scored all poses for each complex in the PDBbind Test Set, taking the highest predicted affinity as the value for that complex, as this was found to be the most effective strategy under cross-validation (Experiments 1.2 and 2.2).

We examined the influence of protein similarity between the training and test sets by excluding from the training set any structure with a protein whose sequence identity was above a given threshold to any protein in the test set. Similarly, we examined the influence of ligand similarity between the training and test sets by excluding from the training set any structure with a ligand whose 2048-bit ECFP4 fingerprint Tanimoto similarity was above a given threshold to any ligand in the test set.

#### Effect of training and testing using docked poses

Figure 1 shows the Pearson correlation coefficient between the predicted and experimental binding affinity achieved by the LB, SB, and HB models on the PDBbind Test Set, when trained and tested using either crystallographic binding poses or docked poses. Figure 1A shows how their performance varies with the maximum permitted protein sequence identity between the training and test sets, and Figure 1B shows how performance varies with the maximum permitted ligand Tanimoto similarity between the training and test sets.

Regardless of the maximum level of protein or ligand similarity permitted between the training and test sets, both the SB and HB models perform worse when trained and tested on docked poses (dotted lines) than when trained and tested on crystallographic poses (solid lines; in Figure 1, the dotted lines are always below the corresponding solid lines for the SB and HB models). The LB model performs identically when using docked and crystallographic poses: the RDKit molecular descriptors used by the model are independent of the pose of the ligand, so the model is unaffected by the use of different poses. The LB model actually performs better than the SB model using docked poses when no data are excluded from the training set. The LB model's performance drops below that of the SB model when complexes with 100% protein sequence identity to those in the test set (Figure 1A), or with a ligand whose Tanimoto similarity was greater than 0.6 to those in the test set (Figure 1B), are excluded from the training and test sets, the LB model always achieves a Pearson correlation coefficient greater than 0.55, comparable to the performance of many classical scoring functions,<sup>4-6</sup> indicating that these ligand-based features capture useful information for affinity prediction.

The HB model, which combines structure-based and ligand-based features, consistently outperforms the SB and LB models when using docked poses (in Figure 1A and 1B, the dotted yellow line is always above the dotted red line and blue line). Furthermore, the HB model trained and tested on docked poses has comparable performance to the SB model when trained and tested on crystal poses (in Figure 1A, the dotted yellow line is above the solid red line; in Figure 1B, the dotted yellow line is nearly identical to the solid red line for a Tanimoto similarity threshold greater than 0.5). Furthermore, the HB model is less deleteriously affected by the use of docked poses in place of crystal poses than the SB model (see, in Figure 1, the dotted yellow line much closer to the solid yellow line than the dotted red line is to the solid red line). This suggests not only that combining ligandbased and structure-based features leads to more accurate binding affinity predictions, but that the addition of ligand-based features to a structure-based scoring function can help to compensate for the errors in affinity prediction that may result from the use of potentially inaccurate docked poses.

All three models (LB, SB, and HB) are strongly affected by the similarity between the training and test sets, with the exclusion of training set complexes with similar proteins or ligands to those in the test set significantly reducing performance. These results echo our earlier results<sup>15</sup> and those of Su *et al.*<sup>33</sup>, indicating that even when potentially less-accurate binding poses are used, it is necessary to consider the effect of biases in the available structural data when training and evaluating models. The inclusion of ligand-based features in structure-based models always improves performance when using docked poses, and only ceases to improve performance when using crystal poses if the maximum fingerprint Tanimoto similarity between ligands in the training and test set is less than or equal to 0.5. This suggests that the inclusion of ligand-based features is a robust method of enhancing scoring function performance, particularly when using docked poses in both training and testing.

#### Effect of training using multiple docked poses

We also checked whether training on multiple docked poses for each ligand affected performance on the PDBbind Test Set. For this we focused on the HB model as it consistently outperformed the LB and SB models in our previous experiments. We repeated the above experiment for the HB model, this time using all of the docked poses we generated for each ligand in the PDBbind Training Set, as described earlier. In contrast with the cross-



Figure 1: Pearson correlation coefficient between predicted and experimental pK values on the PDBbind Test Set, for varying levels of protein sequence identity or ligand fingerprint similarity permitted between the training and test set. Solid lines show performance when trained and tested using crystallographic binding poses; dotted lined show performance when trained and tested using docked poses. The maximum permitted level of protein sequence identity (A) or ligand fingerprint Tanimoto similarity (B) between the training and test set is shown along the horizontal axis. Note: both (A) and (B) have the same vertical axis.

validation experiment, training on multiple docked poses for each ligand significantly reduced the performance of the HB model on the PDBbind Test Set (Supporting Information Figure 4). To understand how training on multiple poses for each ligand affects the HB model, we examined the relative importance of the features used by the RF. We found that when multiple poses for each ligand are used for training, ligand-based features become more important than when only a single pose is used in training (Supporting Information Figure 5), suggesting that using multiple poses for each ligand introduces additional noise into the structure-based features that causes the RF to rely more on ligand-based features.

#### Generalising to unseen proteins remains challenging

#### Validation on PDBbind target clusters

To understand how the HB model might be expected to perform on an unseen target using docked poses, we clustered the PDBbind Training Set at 90% protein sequence identity and identified the six largest clusters. These correspond to the proteins HIV-1 protease, carbonic anhydrase, trypsin, thrombin, heat shock protein  $90\alpha$  (HSP $90\alpha$ ), and coagulation factor X. We also identified all proteins in the PDBbind Training Set that were unique at the 90% sequence identity threshold and grouped these into a seventh cluster of 'singletons', forming a highly diverse set of proteins. For each cluster, we trained the HB model on the remaining complexes in the PDBbind Training Set, using either the pose ranked highest by Smina. We focused on the HB model as it consistently outperformed the LB and SB models in our previous experiments.

The results of this experiment are shown in Table 4. Overall, performance varies greatly across the seven clusters and, with the exception of trypsin ( $\rho_p = 0.746$ ) and HSP90 $\alpha$ ( $\rho_p = 0.591$ ), the correlation between predicted and experimental affinity achieved by each model is poor. These results are similar to those of Kramer and Gedeck,<sup>34</sup> who showed that the performance of RF-Score<sup>35</sup> varies greatly when validated on held-out clusters of PDBbind data, with worse overall performance than when validated on the PDBbind 2007 Core Set. This suggests that performance on diverse benchmark sets such as the CASF test sets is not necessarily indicative of a scoring function's ability to accurately rank ligands of a single target within that benchmark when that target is held out of the training set.

Table 4: Pearson correlation coefficients achieved by the HB model on seven clusters of the PDBbind Training Set when trained on the remaining structures in the PDBbind Training Set. For the training set, the pose ranked highest by Smina was used. For the test sets, all docked poses were scored and the highest value taken as the predicted binding affinity for that ligand.

Cluster	$ ho_{ m P}$
HIV-1 Protease	0.450
Carbonic Anhydrase 2	0.460
Trypsin	0.746
Thrombin	0.316
$HSP90\alpha$	0.591
Coagualtion Factor X	-0.101
Singletons	0.500

#### Validation on the Updated DUD-E Diverse Subset

Having found that the performance of the HB model varies greatly between different protein targets in the PDBbind Test Set, we next investigated how well the model generalised to novel binding data from outside the PDBbind database. To do this, we created and used used the Updated DUD-E Diverse Subset of targets: serine/threonine-protein kinase AKT (AKT1), cytochrome P450 3A4 (CP3A4), glucocorticoid receptor (GCR), HIV-1 protease (HIVPR), HIV-1 reverse transcriptase (HIVRT), and kinesin-like protein 1 (KIF11).

Table 5 shows the Pearson correlation coefficients achieved by the HB model for each target when trained on the PDBbind Training Set, and under both intra-target and intertarget validation, as described in the Methods section. In each case, we trained on the pose ranked top by Smina and tested by scoring all diverse docked poses and taking the highest predicted binding affinity for each ligand.

Using the PDBbind Training Set the performance for all six targets is poor, with the HB model achieving a Pearson correlation coefficient below 0.5 for each target. These results are in stark contrast with the strong performance on docked poses of the PDBbind Combined

Core Set, indicating that even under strict training and validation conditions, the model generalises poorly to data sourced from outside the PDBbind database. One possible source of this difference in performance is that, as previously discussed, the docked poses for the PDBbind data were generated by re-docking each ligand into the corresponding crystal structure of its protein target, so any induced fit effects are already captured in the structure of the protein. Because of this, the docked poses for the PDBbind data are likely to be more accurate than those for the Updated DUD-E Diverse Set. In addition, any uncertainties in the more recent ChEMBL binding affinity data could also contribute to noise and performance degradation when training and validating the model.

Under intra-target training and validation, performance varies greatly between targets, but overall performance is much better than when the models were trained on PDBbind data. For each target, the HB model achieves a Pearson correlation coefficient,  $\rho_{\rm p}$  in the range of 0.5 to 0.8, in contrast to the weak correlations obtained when training on PDBbind data ( $\rho_{\rm p} < 0.5$ ). This indicates that the HB model is capable of making accurate predictions using docked poses of ligands for a protein target, provided it has previously seen examples of ligands for that target.

In contrast with intra-target validation, under inter-target validation the HB model fails to achieve any meaningful correlation between the predicted and experimental pK values for any of the targets ( $\rho_p < 0.4$ ), indicating that the model is unable to generalise to a previouslyunseen protein target. With the exception of CP3A4, the predicted pK values also span a much narrower range of values than the experimentally-determined values (Supporting Information Figures 10 and 11), suggesting that under this training regime, the model is simply unable to differentiate reliably between different ligands for the unseen target. Similar results were obtained using the LB and SB models, indicating that both structure-based and ligand-based RF models will struggle to generalise to novel targets.

Finally, we repeated the inter-target and intra-target validation experiments, adding the PDBbind Training Set to the Updated DUD-E Diverse Subset data, and found that there was no improvement in performance when combining these two sources of training data. This indicates that a larger, more diverse training set does not necessarily translate into improved performance; we made a similar observation in previous work<sup>15</sup> when we showed that the improved performance of several scoring functions on the CASF test sets when training on the PDBbind General Set instead of the PDBbind Refined Set could be attributed not to the increased size and diversity of the General Set, but to its increased representation of the protein targets present in the test sets.

One possible explanation for these results is that docking the ligands for each DUD-E target into a single protein structure provided by DUD-E for that target does not adequately account for induced fit effects, in contrast with docking a ligand into its cognate protein structure and concomitant induced fit, such as when re-docking the PDBbind complexes. This would result in the protein-ligand interaction features failing to capture important interactions between the protein and the ligand, as an accurate docked pose with respect to the correct conformation of the protein's binding pocket is not available. This highlights the need for new experiments to look at the influence of induced fit on docking and subsequent binding affinity predictions when using machine learning models.

Table 5: Pearson correlation coefficients achieved by the HB model between predicted and experimental pK of ligands for six protein targets in the Updated DUD-E Diverse Subset. Three different validation regimes were used. Under 'PDBbind Training', each model was trained on the PDBbind Training Set and tested on all ligands for each target. Under 'Intratarget validation', for each target 80% of the ligands were randomly selected to be used as a training set, and the model tested on the remaining 20%. Under 'Inter-target validation', for each target in turn, all ligands for that target were held out as a test set, with the model trained on all ligands for the remaining five targets. The best performance for each target is underlined.

Target	PDBbind	Intra-target	Inter-target
	Training	Training	Training
AKT1	0.418	0.626	0.220
CP3A4	0.320	0.526	0.302
GCR	0.317	0.809	0.218
HIVPR	0.456	0.739	0.135
HIVRT	-0.101	0.574	0.056
KIF11	-0.177	0.743	-0.038

## Conclusions

We have investigated how the use of docked poses in place of X-ray crystallographic binding poses of ligands affects the ability of Random Forest-based scoring functions to predict protein-ligand binding affinity; how best to make use of docked poses when multiple poses are available; and how scoring functions trained on docked poses generalise to novel sets of ligands.

Unlike Li *et al.*<sup>14</sup>, we found that the use of Smina docked poses in place of X-ray crystallographic binding modes for training and validation reduces the performance of a structure-based scoring function. However, a hybrid model that combines structure-based and ligand-based features is less deleteriously affected by the use of docked poses in training and validation than a purely structure-based model. Furthermore, hybrid models trained on docked poses can achieve binding affinity prediction performance comparable to that of a structure-based model that was trained and validated using crystallographic binding modes. Excluding proteins and ligands from the training set that are similar to those in the test set negatively affected scoring function performance, but ligand-based features still improved binding affinity predictions when removing test-set-similar data from the training set, in agreement with our previous work using crystallographic binding. poses<sup>15</sup>

We also investigated the effect of training on multiple diverse docked poses. Under crossvalidation on the PDBbind Refined Set, there was little difference between training on a single pose per ligand and training on multiple poses. However, when we trained our models using multiple poses for each ligand in the PDBbind Training Set and tested on our heldout PDBbind Test Set, performance was substantially worse than when training using a single docked pose for each ligand. To understand how training on multiple poses per ligand affected the model, we examined the relative importance of the features in each model, and found that when both structure-based and ligand-based features are used, ligand-based features become more important when training on multiple poses, suggesting errors in the structure-based features from the additional docked poses degrade their relative utility. Using binding affinity data obtained from the ChEMBL database (version 25) for six of the eight protein targets in the DUD-E Diverse Subset, we investigated how our models performed on unseen data. Under an intra-target validation where a random selection of 80% of the data for a target was used to train the model and 20% of the data was held out as a test set, our model achieved positive Pearson correlation coefficients between the predicted and experimental binding affinites ranging from 0.541 to 0.810, indicating that our hybrid model is capable of accurate predictions on the data obtained from ChEMBL when trained in a target-specific manner. However, under an inter-target validation scenario, where data for five of the six targets were used to train the model, and the remaining target's data held out as a test set, our hybrid model failed to achieve any meaningful correlation between predicted and experimental binding affinity. Combining the ChEMBL data with PDBbind data to form a larger, more diverse training set did not improve performance.

Overall, our results indicate that, on a benchmark such as the PDBbind 2007, 2013, and 2016 Core Sets, the use of docked poses for training and validation decreases the performance of a structure-based scoring function relative to those trained and tested on X-ray crystal structures, in contrast with the findings of Li *et al.*.<sup>14</sup> Similarities between proteins and ligands in the training and test sets has a strong influence on scoring function performance, in agreement with our previous work<sup>15</sup> and recent work by Su *et al.*.<sup>33</sup> Our results once again suggest that the inclusion of ligand-based features in the scoring function helps to counteract this effect. However, our results also suggest that a model trained on PDBbind data tends to generalise poorly to external data sets, indicating that additional validation and benchmarking sets are needed for scoring function development, although this will depend on the level of similarity between the PDBbind complexes and the unseen proteins or ligand. Overall, our results show that, despite potential problems generalising to novel proteins and ligands, the inclusion of ligand-based features in a structure-based scoring function can help to compensate for errors in binding affinity prediction due to the use of docked poses. By opening the door to the use of docked poses, our hybrid approach expands the utility

of machine learning scoring functions for the discovery of novel small molecules for novel targets.

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