Cosolvent-Based Protein Pharmacophore for Ligand Enrichment in Virtual Screening

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ABSTRACT: Virtual screening of large compound databases, looking for potential ligands of a target protein, is a major tool in computer-aided drug discovery. Throughout the years, different techniques such as similarity searching, pharmacophore matching, or molecular docking have been applied with the aim of finding hit compounds showing appreciable affinity. Molecular dynamics simulations in mixed solvents have been shown to identify hot spots relevant for protein–drug interaction, and implementations based on this knowledge were developed to improve pharmacophore matching of small molecules, binding free-energy estimations, and docking performance in terms of pose prediction. Here, we proved in a retrospective manner that cosolvent-derived pharmacophores from molecular dynamics (solvent sites) improve the performance of docking-based virtual screening campaigns. We applied a biased docking scheme based on solvent sites to nine relevant target proteins that have a set of known ligands or actives and compounds that are, presumably, nonbinders (decoys). Our results show improvement in virtual screening performance compared to traditional docking programs both at a global level, with up to 35% increase in areas under the receiver operating characteristic curve, and in early stages, with up to a 7-fold increase in enrichment factors at 1%. However, the improvement in pose prediction of actives was less profound. The presented application makes use of the AutoDock Bias method and is the only cosolvent-derived pharmacophore technique that employs its knowledge both in the ligand conformational search algorithm and the final affinity scoring for virtual screening purposes.

INTRODUCTION

Drug discovery is an expensive endeavor, costing between 0.16 and 1.8 billion dollars to introduce a new drug to the market. The need for new compounds and more efficient tools to obtain them is illustrated by the outbreak of infectious diseases, boosted by the emergence of drug resistance in microorganisms such as Mycobacterium tuberculosis, Staphylococcus aureus, or Klebsiella pneumoniae among others and the need for better treatment of neglected diseases such as malaria, dengue, or sleeping sickness. Furthermore, the discovery of new molecular targets to treat complex diseases such as cancer, autoimmune, and neurodegenerative diseases also requires the development of new drug candidates and pharmacological tools to elucidate their roles in physiological and pathological contexts.

Virtual screening (VS) of large compound databases, looking for potential ligands of a target protein, is a major tool in computer-aided drug discovery. Its main purpose is to find hit compounds with a certain affinity, usually around 1 μM, which can then be further optimized to become lead compounds and drug candidates faster, more efficiently, and at lower cost than with experimental high-throughput screening campaigns.

Briefly, VS consists in rapidly estimating the binding free-energy of compounds to a given target by computational means and choosing a subset of these compounds with higher predicted affinities to be tested experimentally. A widespread VS technique starts from the target protein structure in the so-called docking methods and explicitly studies the interaction between the protein and the compounds to be evaluated.

Docking-based VS is far from perfect, and there are several issues that require improvement. Correct pose prediction is still a difficult task and accurate free-energy estimations are a major challenge, in part due to ignoring the flexibility of the target and desolvation effects. Additional information about the target of interest, for example, ligand interactions revealed in a large number of cocrystal structures or similarity patterns between known ligands, is sometimes available, and its incorporation into VS campaigns can increase success rates. The improved performance of knowledge-based docking is...
DOCK613 has a conformational search option to bias the perform template guided (or tethered) docking. Finally, thus bias the results toward important interactions and also to constrain generated poses to satisfy pharmacophores and information from water and solvent sites. We have previously shown applications of solvent sites to protein surface where the probability of finding water or solvent sites, that is, conformed regions close to the hot spots. Molecular dynamics (MD) simulations of a protein in explicit water or mixed solvents that mimic specific types of ligand interactions have been increasingly used to analyze protein binding sites and to reveal protein–ligand interaction hot spots. MD allows studying the structure and dynamics of water and cosolvent molecules which, as a consequence of the shape and charge distribution of the protein surface, are arranged inhomogeneously in the solvation shell. This gives rise to solvent sites, that is, confined space regions close to the protein surface where the probability of finding water or cosolvent molecules is significantly higher than that of the bulk. We have previously shown applications of solvent sites to binding mode and binding free-energy predictions, especially using ethanol as a probe that is able to capture in a single molecule the hydrophobic, hydrogen bond donor and hydrophilic and hydrophobic ethanol sites that are common in drug-like molecules. On the basis of our work on ligand docking using information from water and solvent sites, we have recently developed AutoDock Bias for performing biased docking with AutoDock422 program. The versatile definition of the different types of biases in AutoDock Bias allows guided docking toward pharmacophoric interactions in a straightforward way for hydrogen bond and hydrophobic/aromatic interactions, and precise localization of any desired atom (e.g., metal) or group (e.g., substructure core of a congeneric ligand series or for fragment growth) in a defined region of the target protein, which may also be used as an anchor for covalent docking studies. In the present work, we have retrospectively tested the performance of AutoDock Bias using the information from MD-derived solvent sites for its application in VS campaigns. As previously stated, we chose ethanol as a cosolvent, and once the ethanol sites were computed from the corresponding MD simulations (see Computational Methods), we implemented a biased docking scheme based on their location, binding free energy, and dispersion. We selected nine relevant target proteins that have a set of known ligands or actives and compounds that, presumably, do not bind to the protein ( decoys). Our results show that supplementing the docking conformational search algorithm and scoring function for each target yielded improved VS performance both globally, as seen in receiver operating characteristic (ROC) curves and area under the ROC curve (AUC), and in early enrichment, shown by semilogarithmic ROC curves, log AUC, EF1%, EF10%, and Boltzmann-enhanced discrimination of ROC (BEDROC) metrics.

## RESULTS AND DISCUSSION

The results are organized as follows. We first present the VS performance analysis of our biased docking method, AutoDock Bias, using MD-derived solvent sites and compare it to conventional docking programs (AutoDock4 and Vina) on nine different protein targets. Then, we move on to evaluate the relative scoring and pose prediction of known active molecules by the different docking methods to gain insight into the origin of observed improvement with AutoDock Bias. Finally, we outline the limitations of the proposed method and compare it with similar approaches.

For each of the nine proteins, we ran a set of three 20 ns MD simulations in an aqueous solution of ethanol 20% v/v and compute the ethanol interaction sites (see Computational Methods). Interestingly, and confirming our previous study, these MD-derived ethanol sites match the known pharmacophoric ligand interactions for these systems to a great extent. Details for each protein are included in the Supporting Information, and a graphical description is shown in Figure S1, where ethanol sites are superimposed on representative ligands. For their application in AutoDock Bias, both hydrophilic and hydrophobic ethanol sites were identified on each target binding/active site, ranging from one to five

![Figure 1](image-url)
interaction sites of each kind. Hydrophilic sites were used to bias hydrogen-bond interactions according to their donor and/or acceptor capacity, and hydrophobic sites were used to bias hydrophobic interactions involving aromatic rings (see Computational Methods for details).

We tested the high-throughput performance of AutoDock Bias in VS campaigns in a retrospective manner. Seven of the nine studied systems are relevant target proteins taken from the DUD-E database.24 The two remaining systems, BRD4 and galectin-3, were both subjected to the DUD-E protocol to generate decoy molecules (http://dude.docking.org/generate)24 and then added to the original set to enrich its diversity. All targets, therefore, have a set of (a) known true ligands called "actives" and (b) decoys or compounds that have similar physical and chemical properties as the actives but, presumably, do not bind to the target protein. After docking actives and decoys to their target, a ranking of the compounds was built based on their score, and ROC-derived measures such as AUC, log AUC, EF1%, EF10%, and BEDROC were calculated as defined in the Supporting Methods. The whole process is outlined in Figure 1.

**Figure 2.** ROC curves for the docking of actives and decoys against nine protein targets. Results obtained with standard AutoDock4 (blue) and Vina (yellow) are compared with those obtained by AutoDock Bias using ethanol hydrophilic sites (red) and both ethanol hydrophilic and hydrophobic sites (green). The gray straight line is expected for a random selection of compounds from the database. The informed p-values correspond to the hypothesis test about the difference in AUC between the conventional method (AutoDock4) and AutoDock Bias (using ethanol hydrophilic sites in red and ethanol hydrophilic and hydrophobic sites in green).

**AutoDock Bias for VS.** To quantify the ability of AutoDock Bias and the standard docking methods to rank active compounds in the first positions, that is, their VS performance, we calculated different metrics that evaluate both the overall performance (ROC curve and AUC) and the early ligand enrichment (semilogarithmic ROC curve, log AUC, enrichment factors, and BEDROC). The early ligand enrichment is of particular interest because, in practice, only a small subset of the original database, usually less than 10%, is experimentally tested. Therefore, it is desired that the first 1 or 10% of the ranking is significantly enriched in true ligands that can move on to the next stages of the drug-development process.
Figures 2 and S2 show the ROC and semilogarithmic ROC curves, respectively, for nine protein targets obtained with different docking methods, two standard docking programs (AutoDock4 in blue and Vina in yellow) and AutoDock Bias using ethanol sites. For AutoDock Bias, the performance using only hydrophilic ethanol sites and both hydrophilic and hydrophobic ethanol sites is shown in red and green, respectively. The expected behavior for a random extraction of compounds is also depicted in gray for comparative purposes. Table 1 shows the results obtained for ROC-derived metrics corresponding to the same set of proteins and docking methods.

The first general observation emerging from Figures 2 and S2 and Table 1 is that conventional methods, AutoDock4 and Vina, have very different performances depending on the tested system, showing poor metrics for several of them. This occurs both globally, as represented by ROC curves rising only slightly above the line corresponding to a random extraction of compounds from the database and AUC values below 0.70, and also at an early stage, observed in semilogarithmic ROC curves not far from the random curve and log AUC values below 0.15. These arbitrary thresholds were derived from the mean values for each metric considering conventional methods. On three of the studied target systems, namely Factor Xa, DHFR, and CDK2, conventional methods have good (AUC > 0.70, log AUC > 0.15) to excellent (AUC > 0.80, log AUC > 0.25) performance, whereas on the other six, poor metrics are obtained for either one or both methods. Interestingly, both methods fail for the two systems that were not retrieved from the DUD-E database (BRD4 and galectin-3). The case of galectin-3 is particularly striking since AutoDock4 shows a worse overall ranking than what would be obtained by random selection, whereas the early enrichment measured by log AUC is practically zero. This is in agreement with the results shown in Table 1.

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aMean (standard deviation) for all systems. bValues highlighted in bold represent significant improvement in AUC and/or log AUC over standard docking procedures (p < 0.05, see text).
with previous observations of poor results for carbohydrate docking. Overall, the comparative performance between both conventional methods, especially at early enrichment, shows that for most systems, AutoDock4 gives substantially better results than Vina (AmpC, BRD4, CDK2, Factor Xa, FGFr1), whereas for some others, the trend is reversed (DHFR, galectin-3, HIV protease). Statistical differences between AutoDock4 and Vina performances are shown in Table S1, and a discussion in the context of a recent study that concluded that Vina outperforms AutoDock4 using the CASF2013 data set is presented in the Supporting Discussion.

On the contrary, AutoDock Bias achieves good performance (AUC > 0.70, log AUC > 0.15) in five systems and excellent performance (AUC > 0.80, log AUC > 0.25) in other three systems. The only target where the performance of the biased methods is still poor, albeit better than the conventional ones, is phosphodiesterase 5A. Also, when using only ethanol hydrophilic (ethanol −OH) sites as a bias, a low early enrichment was achieved for FGFr1 and HIV protease. In relative terms, both biased methods have superior performance than conventional ones for all systems except FGFr1 at early enrichment. The conformation of the active site of this flexible kinase target is fixed in the DUD-E structure to the so-called DFG-in state, which was the one sampled during the MD simulations used to determine the ethanol sites. It seems that the bias toward interactions in this particular conformation strongly penalized DFG-out ligands (actives) that might be present in the DUD-E dataset. This is expected since the DFG-in and DFG-out conformations are structurally and energetically dissimilar. Finally, considering all targets, an increase in mean results is obtained for all metrics with the biased methods, although this has a limited value especially due to the difference in performance between systems. In this regard, the lower standard deviation values highlight that AutoDock Bias performance using both hydrophilic (−OH) and hydrophobic (−CH3) ethanol sites is more consistent between diverse systems than conventional AutoDock4.

To statistically analyze the observed differences between conventional methods and AutoDock Bias, a one-tailed hypothesis test was performed using bootstrap (see Supporting Methods). The obtained p-values for the AUC are shown next to the ROC curves (Figure 2) and for the log AUC next to the semilogarithmic ROC curves (Figure S2) from which they are derived, for both biased methods. Taking the typical value of 0.05 as the significance level for the test, the AUC and log AUC values obtained using AutoDock Bias with both hydrophilic and hydrophobic ethanol sites are significantly higher than those of conventional AutoDock4 for seven out of the nine systems. The systems where no significant improvement (nor decline) was observed are FGFr1/AmpC for AUC and FGFr1/FXa for log AUC. A similar enhancement is achieved by the biased method with only ethanol −OH sites for the AUC values, but the improved performance is reduced to six out of nine targets for log AUC. The tests were performed against AutoDock4 because it is the conventional method that gave the best overall results and the one used as a baseline for our energy modifications. Beyond the statistical analysis of AUC and log AUC values, there are five targets (AmpC, BRD4, DHFR, galectin-3, and HIV protease) for which the improvement with AutoDock Bias using both types of ethanol sites is evident both qualitatively, by the separation between the corresponding ROC and semilogarithmic ROC curves, and quantitatively, especially in terms of very early enrichments: EF1% is 2–7 times higher than those obtained by the conventional methods. For example, AmpC has ca. 7-fold increase in its EF1% for AutoDock Bias. This means that if a database is composed of 100 000 compounds out of which 100 are true ligands, we expect to experimentally obtain 2 hits if we select compounds based on standard docking software, but instead to get 15 hits if using AutoDock Bias, for every 999 confirmed nonbinders.

To summarize the overall performance of AutoDock Bias using hydrophilic and hydrophobic ethanol sites compared to that of conventional methods, regarding both global and early enrichment, Figure 3 shows their corresponding difference in AUC and BEDROC for all systems. The Δ(AUC) versus Δ(BEDROC) plot displays improvement for AutoDock Bias in practically all of the systems, as revealed by the points in the (+,+ ) quadrant. Most systems have BEDROC differences greater than 0.10, which, according to the statistical interpretation of this parameter (see Supporting Methods), are values of substantial improvement. As previously analyzed, the only case in which AutoDock Bias does not attain better early parameters than AutoDock4 is FGFr1, yielding a value of BEDROC lower in 0.02 units and a drop of 5.7% in EF1%. However, the loss of enrichment is quickly reversed at EF10% and thereafter, and improvement is observed with respect to Vina at all stages. Finally, the correlation for the increase in both global and early metrics seen in Figure 3 indicates that a general improvement is achieved with the biased method. However, the rise in global enrichment appears somewhat more moderate than the increase in early enrichment, suggesting that the bias causes a substantial increase in the score of some actives relative to decoys, but cannot fix grossly incorrect predictions, for example, due to inadequate protein conformation.

**Scoring and Binding Mode of Actives.** To analyze the effect of the biased conformational search and scoring on the docking of known binders, we compared it to conventional docking methods regarding (a) the relative ranking of active molecules and (b) the binding mode prediction of actives with a known bound crystal structure (see Computational Methods).

The rankings of active molecules obtained with standard AutoDock4 and AutoDock Bias using both hydrophilic and hydrophobic ethanol sites were compared by calculating the

![Figure 3. Δ(AUC) vs. Δ(BEDROC) between AutoDock Bias using ethanol hydrophilic + hydrophobic sites and AutoDock4 (dots) or Vina (crosses). Results are color coded according to each system.](image-url)
Kendall tau coefficients (Table S2). Eight out of nine systems display a correlated order of active molecules in both rankings ($\tau \geq 0.5$). Therefore, when using AutoDock Bias, a general rise in the ranking of actives as a group with respect to decoys is accomplished, whereas the order between actives themselves is not significantly altered when compared to conventional docking results. This highlights that the biased docking method maintains the relative binding affinity predicted for actual ligands by AutoDock4, whose scoring function was calibrated with known binders, while better discriminating between binders and nonbinders by enforcing specific target important interactions.

We also compared actives docking pose with their crystallized structure with the aim of discerning if better ranking correlates with a more accurate pose prediction. Heavy atom root-mean-square deviations (rmsds) between the predicted pose for active molecules and the co-crystallized conformation of the same compound or a similar one (Tanimoto similarity index $\geq 0.85$) are presented in Figure 4. In five of seven systems, there was an improvement in pose prediction using AutoDock Bias: mild (5% more actives with rmsd $< 2 \text{ Å}$) for PDE5A and DHFR, moderate (10%) for BRD4, and significant (45%) for FGFr1 and AmpC. However, there is slightly worse prediction for CDK2 (−5%) and a moderate one (−16%) for Factor Xa. Therefore, although improving pose prediction in the majority of tested proteins, AutoDock Bias performance remains system-dependent for significantly improving binding mode calculations.

It can be seen from Figure 4 that there are several systems with highly inaccurate binding mode predictions (rmsd $> 4 \text{ Å}$). In BRD4, the reason seems to be that the main driving force for the binding is a single hydrogen bond interaction with Asn140, aided by variable nondirectional hydrophobic interactions. An additional difficulty for the docking is that there are known interactions bridged by water molecules. The poor pose prediction holds even with the biased method and is related to the lack of solvent sites—except the ones near the Asn140—that would aid in positioning the ligands correctly in the whole binding site. On the contrary, the enhanced formation of the hydrogen bond between Asn140 and the ligands in the biased docking runs is enough to improve the VS performance, as revealed by significant increases in AUC and log AUC (Table 1). Unlike BRD4, DHFR constitutes a case where improvement in VS performance by the biased method is explained by better binding mode prediction. The conventional method has problems with several similar active ligands that repeatedly appear with their terminal fused aromatic rings inverted in the binding site, dramatically changing the interaction pattern (Figure S6A,B). The bias toward the hydrogen bond interaction between Glu30 and the diamino-pyrimidine ring (Figure S1D) corrects this issue for the biased method. Last, there are problematic ligands whose poses cannot be predicted by any of the methods and correspond to structures of considerable size with around 14 torsions and 50 atoms excluding nonpolar hydrogens (Figure S6C). Finally, for PDE5A and FGFr1, there is a pose prediction improvement with the biased method, but as previously discussed, the flexibility of both protein targets prevents from achieving better predictions since the protein conformations from the DUD-E set are substantially different from the holo structures found for certain types of ligands from the active set. Interestingly, the binding mode of ligands is certainly better captured by the biased method for FGFr1, a system whose improvement in enrichment parameters is quite poor (Table 1). We can rationalize this result taking into account that all ligands with rmsd below 2 Å are bound to the DFG-in protein conformation in the corresponding crystal structure, which matches the conformation selected from the DUD-E database for the docking experiments. This points out again that AutoDock Bias is probably incapable of assigning high scores to ligands that bind the DFG-out conformation and are present as actives in the test set.

Finally, it is interesting to dig deeper into the results for Factor Xa that show worse pose predictions with AutoDock Bias than with standard AutoDock4. This observation can be explained by the lack of directionality of the hydrophobic/aromatic pharmacophore derived from the MD, which is the binding driving force for this system. This feature is unique to Factor Xa since all other systems from the set have a robust polar pharmacophore which provides directionality for ligands (Figure S1). In this sense, it should be interesting to derive a better bias for the aromatic moieties that also includes directionality, for example, by using phenol as a cosolvent. In conclusion, the improvement observed in the classification between actives and decoys by the addition of the hydrophobic bias is relevant for VS but needs additional adjustment since ligand correct orientation is not well achieved when no hydrogen bonding is included. As a final remark, several FXa ligands bind benzamidine or other positively charged groups. Ethanol being a neutral molecule was able to identify this pharmacophoric hydrogen bond, but an ionic probe would probably assign better energy prioritization and predict correct poses for more ligands.

Discussion in Relation to Other Methods. Pharmacophores obtained from the protein structure, with no previous ligand-based knowledge, are versatile for the discovery of new drugs since they can be employed in novel targets for which limited information is available. Likewise, in the VS of large amounts of compounds, they increase the probability that the obtained hits have different chemical functionalities than the ligands already tested for the target of interest. Moreover, the
protein-derived pharmacophores obtained in this study consider desolvation in an explicit way and protein flexibility to a certain degree because they were obtained from unrestrained MD simulations in mixed solvents.

To achieve more profound improvements with the developed method, we must answer two central questions. The first one is why some actives still do not appear in the upper part of the ranking. A careful analysis of ligand properties show that, in general, actives that are not retrieved at early or middle stages are relatively small molecules compared to those ranked higher (Figure S4). Although this is expected given that docking score scales with the size of compounds, AutoDock Bias seems less affected by the ligand size effect on the ranking, as revealed by the Spearman’s correlation coefficients closer to 0, than the conventional method for seven of the nine systems (compare Figures S3 and S4). The second question is why some decoys remain in the upper portion of the ranking even when the biased method is applied. We found that many highly ranked decoys adopt excessively folded and seemingly entropically unfavorable conformations that allow them to coincide with several solvent sites and thus increase their estimated binding energy. This is possible because the scoring function has a poor balance between the interaction energy and conformational entropy. On the other hand, we also observed that there are cases in which the best-ranked decoys preferentially replace solvent sites that are not coincident with ligand-derived pharmacophoric sites when compared to best-ranked actives. In general, potential areas for future improvement include the use of additional solvent probes that capture other interaction types (e.g., ionic groups) and taking advantage of the information about displaceable/non-displaceable water molecules that already comes out of the MD simulation in mixed solvents.

Finally, it is interesting to compare our method with other works using MD in mixed solvents in the context of VS. MacKerell’s group pioneered the use of pharmacophore models obtained by cosolvent MD for screening a large number of compounds. In their approach, ligand features are required to match all MD-derived pharmacophoric sites, and the score arises from the rmsd between these paired features, supplemented with free-energy grid calculations for the ligand pose(s). The method is useful to filter compounds that are not able to fulfill the pharmacophoric sites, but does not aid in the conformational search itself. Indeed, it relies on predefined ligand conformations that are not adjusted to the target structure. Nonetheless, since there is no rigid receptor, protein flexibility from MD simulations is totally accounted in the pharmacophoric sites, and excluded space regions are determined from the absence of probe localization during the MD rather than from a given protein surface. On the other hand, Uehara and Tanaka used cosolvent MD to generate an ensemble of protein conformations for docking-based VS. The ensemble selection relies on clustering the conformations according to binding site residues and ranking the clusters by predocking known actives, considering the ones achieving higher scores. Finally, ensemble docking is performed using standard Vina. This approach is totally complementary with our method since the final ensemble docking could be done with AutoDock Bias using solvent sites extracted from the same MD simulations required to obtain the target receptor ensemble. To our knowledge, the AutoDock Bias implementation presented here is the only method where cosolvent-derived information is used to modify the scoring function of a docking program, both for conformational search and final affinity estimation, with the aim of improving VS results.

CONCLUSIONS

In the present work, we demonstrate the potential of biased docking using pharmacophores derived from MD in mixed solvents (solvent sites) to increase the enrichment of true ligands in VS campaigns. Significantly higher AUCs and log AUCs were obtained with respect to conventional docking methods, indicating improvement both in the global ranking of compounds and, especially, in early stages. The latter important issue is additionally reflected in ligand enrichments at 1% which were multiplied by up to 7 times. The most successful variant is to perform MD simulations in the ethanol-water mixture, obtain both ethanol hydrophilic and hydrophobic interaction sites, and add this solvent information to the docking protocol of AutoDock Bias.

COMPUTATIONAL METHODS

Protein and Ligand Dataset. Nine protein targets were selected (Table 2) to test the performance of the biased docking method for VS. They belong to different families, have various functions, and represent a diverse set of targets relating its protein–ligand interactions (Figure S1). The protein structures for seven of the targets were obtained from the DUD-E database, a repository of relevant target protein structures, each bearing a set of true ligands or actives with an experimentally measured affinity below 1 μM and nonbinders or decoys, which are compounds with similar physicochemical properties as the actives but a different topology based on two-dimensional (2D) fingerprints. In addition to these focused...
decays, molecules proved to be experimentally inactive (affinity greater than 30 \( \mu \)M) were also added to the database when that information was available. The DUD-E, together with its predecessor DUD,\(^{31}\) has been proven useful for benchmarking VS procedures.\(^{32}\) The two remaining systems not present in the benchmarking database are BRD4, a promising target for disrupting protein–protein interactions, and galectin-3, a docking-challenging carbohydrate binding protein with a binding site quite exposed to the solvent and known ligands with increased complexity due to high number of rotatable bonds. The structures of these two targets were obtained from the PDB in the holo form with a resolution lower than 2.5 Å and in a conformation that could accommodate the greater number of cocrystallized ligands. These resulted in PDB ID 4xc9 for BRD4 and 1kjr for galectin-3. Subsequently, a procedure analogous to that of the DUD-E was carried out. First, actives were recovered from BindingDB,\(^{33}\) BindingMOAD,\(^{34}\) and ChEMBL,\(^{35}\) databases, only considering compounds with experimentally measured \( K_p \) or IC50 lower than 1 \( \mu \)M against the corresponding target. Then, a structural diversity filter was applied with DataWarrior program,\(^{36}\) based on three-dimensional (3D) similarity according to protein-binding pharmacophoric characteristics (Flexosphere similarity descriptor).\(^{37}\) Finally, the decoys were obtained from these actives through the DUD-E web tool (http://dude.docking.org/generate),\(^{38}\) which recovers 50 decoys extracted from ZINC\(^{18}\) for each active protomer in the SMILES format. SMILES for actives and decoys of BRD4 and galectin-3 are available in the Supporting Information. In summary, the number of active ligands per system ranges from 37 to 537, while the number of decoys varies between 1694 and 35 674, so different sizes of databases were analyzed with the method (Table 2).

**Workflow and ROC Metrics.** To evaluate the performance of different docking methods in VS campaigns, we followed the process outlined in Figure 1 for each target. After retrieving actives and decoys, all compounds were docked against the protein structure stated in Table 2 and Supporting Methods, and a ranking was built according to their predicted affinity. The predominance of active ligands in the first positions of the ranking was quantified by the following metrics: ROC curve, AUC, semilogarithmic ROC curve, adjusted area under the ROC curve, enrichment factors at AUC, semilogarithmic ROC curve, and a ranking was built according to their predicted activity. The calculation of the different metrics from the rankings was made with R.\(^{40}\) In particular, we use several functions from the “ROCR” package\(^ {41}\) to calculate ROC curves and AUCs, the “pracma” package to obtain the log AUC through integration using trapezoids, and the “pROC” package\(^ {42}\) to calculate the hypothesis test for the difference in AUC. The hypothesis test for the difference in log AUC was programmed in R following the guidelines included in the Supporting Methods.

**MD in Mixed Solvents.** The MD simulations in aqueous ethanol for AmpC, CDK2, DHFR, Factor Xa, FGF receptor 1, and PDE5A were previously performed and described.\(^ {18}\) BRD4, galectin-3, and HIV protease were simulated following the same protocol. Initial structures were downloaded from the PDB database\(^ {43}\) (PDB IDs 4xc9 for BRD4, 1kjr for galectin-3, and 1xl2 for HIVpr). All nonstructural ions and solvent and ligand molecules were removed. Missing side chains and hydrogen atoms were added using the LEaP module from Amber package (version 16).\(^ {44}\) Standard protonation state at physiological pH was assigned to all ionizable residues, except for HIV protease that was simulated with Asp124 protonated. The structures were then solvated with a truncated octahedral box of ethanol 20%V/V extending at least 10 Å from any protein atom using MMDBx program.\(^ {55}\) TIP3P model was used for water molecules, and forcefield parameters for ethanol were assigned as previously reported.\(^ {18}\) Amber ff14SB forcefield was used for protein residues. All MD simulations were performed with the PMEMD implementation of SANDER for GPU from Amber 16.\(^ {56}\) Details on ethanol concentration and simulation protocol can be found in the Supporting Methods.

**MD-Derived Solvent Sites.** The MD-derived solvent sites (ethanol sites) were obtained as explained in our previous work.\(^ {18}\) Briefly, ethanol sites are defined as space regions adjacent to the protein surface where the probability of finding a probe atom of this cosolvent is higher than that in the bulk. We obtained both types of ethanol sites: hydrophilic sites in regions where the –OH resides for a significant amount of time during the MD simulation and hydrophobic sites in regions where the terminal –CH3 had a considerable occupancy. The preference of the ethanol probe for being in the solvent site with respect to the bulk is given by the free energy of the site defined in eq 1

\[
\Delta G = -RT \ln \frac{N_{\text{solvent site}}}{N_{\text{bulk}}}
\]

where \( N_{\text{solvent site}} \) is the number of times a probe atom is found inside the solvent site during the MD simulation and \( N_{\text{bulk}} \) is the number of times a probe atom is found in the bulk inside a sphere with the same volume of the solvent site (arbitrarily taken as 1 Å³). Besides its free energy, a solvent site is also described by its coordinates, that is, the center of mass of the probe atoms defining the site and its radius or dispersion parameter, that is, the radius necessary to get a sphere containing 90% of the probe atoms defining the site. Ethanol sites were obtained exclusively for the binding (or active) site of each target. For each protein, the binding pocket was defined using all protein–ligand cocrystal structures available in the PDB, having a resolution below 2.5 Å and a reported binding affinity (\( K_d, K_i \) or IC50) below 1 mM. All protein residues that had at least one atom within 4 Å from any ligand atom in at least one structure were considered part of the pocket. To select the cosolvent pharmacophoric points (solvent sites) for the VS application, we first identified all possible solvent sites in the binding pocket. Then, we kept those with \( \Delta G \) of binding above 0.95 kcal/mol (\( \Delta G < -0.95 \) kcal/mol), a threshold representing a probe-finding probability at least 5 times higher than the one in bulk, eq 1). Similar cutoffs have been used in previous studies for deriving pharmacophore profiles.\(^ {5,26}\) Finally, solvent sites clashing with the particular protein structure selected for docking, as a consequence of the receptor flexibility during the MD, were
removed. The distance cutoff between a solvent site and any protein heavy atom below which the solvent was discarded was set at 2.1 Å for hydrophilic sites and 2.9 Å for hydrophobic sites (a 0.5 Å tolerance from the minimum values for hydrogen bonds and aliphatic/aromatic interactions, respectively, reviewed by Bissantz et al.46). As a final remark, the solvent sites overlapping the NADP cofactor from DHFR were also removed because the cofactor was retained for the docking as if it was part of the receptor.

**Conventional and Biased Docking. Protein, Ligand, and Grid Preparation for Docking.** Protein structures were obtained from the DUD-E database for seven systems (AmpC, CDK2, DHFR, Factor Xa, FGF receptor 1, HIV protease, and PDE5A) and directly from the PDB for BRD4 (4zc9) and galectin-3 (1kjr). They were checked and protonated when needed with Maestro45 and then converted to PDBQT with AutoDockTools22 (prepare_receptor4.py). Structural details can be found in the Supporting Methods. Each receptor was kept rigid for docking. Structural ligand files were obtained from the DUD-E database in the SDF format and converted to PDBQT format with Open Babel.48 For the systems not present in the DUD-E database, the 3D structures of the compounds were generated with LigPrep49 from the previously obtained SMILES (see “Protein and Ligand Dataset”), considering stereoisomers, tautomers, and ring conformers.

For building the grid that defined the docking site for each target, all protein—ligand crystallographic complexes available from the PDB, with resolution below 2.5 Å and a reported binding affinity (Kd, Ke, or IC50) below 1 mM, were used. For systems with less information, such as galectin-3, all complexes from the PDB were retrieved. The structures were aligned according to the binding site residues to the protein structure from the PDB. The 3D structures of the ligands were obtained from the DUD-E database, the 3D structures of the libraries (PDBQT) are provided as Supporting Content, and docking grids position and dimensions are available in Table S3.

**AutoDock4 and Vina.** On the one hand, conventional docking experiments were performed with AutoDock422 program (version 4.2.6). Briefly, based on the protein structure, the energy maps were computed as usual inside the grid. One hundred independent docking runs were performed, and the resulting poses were clustered according to the ligand heavy atom rmsd using a cutoff of 2 Å, thus defining a population for each cluster. The Lamarckian genetic algorithm was applied with a maximum of 27 000 generations or 2 500 000 energy evaluations. On the other hand, when using Vina23 for the docking runs, the parameters of the default search algorithm were used, and nine binding modes were generated for each ligand. The energy maps were calculated internally during the docking experiment using the same position and dimensions as for AutoDock4.

**AutoDock Bias with Solvent Sites.** The solvent site-biased docking method has been described in detail in our previous work.18 Here, we only use solvent sites obtained from MD simulations in ethanol as a cosolvent because of the specificity and precision of the probe and the binding mode results previously shown.19 To adapt the ethanol sites obtained from a flexible MD simulation to the rigid structure used in docking experiments, a positional correction for the hydrophilic sites was made according to their hydrogen bond donor and/or acceptor capacity. The coordinates of the hydrophilic ethanol sites were translated to the ideal location of the hydrogen bond formed according to the conformation of the receptor group of the protein, following the guidelines of Boobyer et al.50 on which AutoDock4 scoring function is also based. Thus, acceptor sites remained located at ideal positions for heavy atoms and donor sites at those for H atoms. A single solvent site was chosen for each protein interactor atom, that is, not more than one solvent site of the same type can interact with the same protein atom. Then, AutoDock4 grid maps were modified such that O/N acceptor atoms from the ligand have an energy reward in the corrected location of acceptor ethanol sites while ligand H donor atoms have it in the corrected location of donor ethanol sites. In addition, aromatic ring centers from the ligand have an energy reward in the regions where hydrophilic ethanol sites were established (no positional correction was done because of lack of well-established directionality for these interactions). The rationale for guiding aromatic rings to hydrophobic sites derived from a nonaromatic probe comes from the finding made by Barril’s group,15 which showed that the terminal C of ethanol was able to reproduce ligand aromatic interactions with the same specificity as aliphatic ones. This was further supported by our previous work,18 where we showed that ethanol has a sensitivity around 70% for ligand hydrophobic interactions and that, even though it does not have aromatic properties, has a slight preference for reproducing interactions of ligand aromatic rings compared to aliphatic rings or side chains.

The magnitude of the energy reward applied to the modified maps is directly related to the free energy of the ethanol sites according to eq 2

$$
\Delta G_{\text{bias}} = \Delta G_{\text{ori}} + \sum_{i=1}^{N} \Delta G_{i} \times e^{-\frac{(x-x_{i})^2 + (y-y_{i})^2 + (z-z_{i})^2}{r_{i}^2}} $$

where $\Delta G_{\text{bias}}$ is the resulting modified score, $\Delta G_{\text{ori}}$ is the original AutoDock4 score, $\Delta G_{i}$ is the free energy of the i-th solvent site, $(x,y,z)$ are the grid point coordinates, $(x_{i},y_{i},z_{i})$ and $r_{i}$ are the coordinates and radius of the i-th solvent site, and the sum extends over the total number $N$ of ethanol sites. Modified OA/NA/HD energy maps built by AutoDock Bias are then used in the same way and in place of conventional AutoDock4 maps. For the aromatic bias, a new map was built using eq 2 with $\Delta G_{\text{ori}} = 0$ and affecting a dummy atom type located in the center of ligand aromatic rings. The rest of docking parameters were kept as with conventional AutoDock4 runs for sake of comparison.

**Final Score.** As it is common with AutoDock4, we used a combined scoring scheme. The consistency of a particular solution in repeated and independent docking calculations was evaluated by gathering the different predicted poses according to the rmsd of their atom coordinates, thus defining a population parameter that counts equivalent poses. Then, the scoring criterion used for each docked ligand consisted in retaining the pose with the lowest free energy of binding among those with a population equal or greater than 20% (if the population resulted in less than 20% for all poses, then the lowest global energy was taken). Finally, a ranking of compounds was generated according to the final score obtained in this way. For the same compound, only the protomer, conformer, tautomer, or stereoisomer with the most
favorable energy score was considered after applying the criterion just discussed. For Vina, the better ranked binding mode was merely taken with it associated score.

**Binding Mode Prediction.** Regarding pose prediction assessment, for each target, cocrystallized ligands with a Tanimoto similarity index equal to or above 0.85 against any of the actives of that target were retrieved from the PDB (see Supporting Methods for notes on the selected similarity threshold). The computation of similarity was done directly on the PDB server, using the extended connectivity fingerprint method from ChemAxon JChem. Target proteins were filtered by UniProt ID with sequence identity above 90% from the reference structure selected for docking. For each corresponding pair (docked active pose/crystallized ligand conformation), the protein structures were aligned according to the binding site, the maximum common substructure (MCS) between the ligands was found, and their heavy atom rmsd was computed using the RDKIT.\(^1\) When many MCSs alignments and/or crystallized ligands were available, the reported rmsd is the minimum of them. Crystallized ligands outside the docking grid were left out of the analysis. The procedure was performed for conventional and biased docking results and comprised 19 ligands for AmpC, 86 for BRD4, 92 for CDK2, 19 for DHFR, 84 for Factor Xa, 9 for FGFr1, and 25 for PDESA. Galectin-3 lacks a proper number of cocrystallized ligands similar enough to the active molecules (it has only three crystallized ligands on the PDB which have binding data on ChEMBL, one of the databases used to build the active molecules set), and HIV protease structure from the DUD-E database differs enough from several other PDB entries, making the alignment unfeasible.

**ASSOCIATED CONTENT**

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jcim.9b00371.

MD-derived solvent sites, semilogarithmic ROC curves for the docking of actives and decoys, number of heavy atoms in the active ligand vs. its position in the ranking, backbone rmsd for BRD4, Galectin-3, and HIV protease during the MD simulations, comparison of AUC and logAUC between AutoDock4 and Vina, Kendall tau correlation coefficient for rankings of actives, grid position and size defining the docking region, AutoDock4 and Vina performances, and supporting methods (PDF)

SMILES for actives and decoys of BRD4 and Galectin-3 (ZIP)

PDBQTs for target proteins (ZIP)

The scripts for obtaining solvent sites are available upon request, and AutoDock Bias\(^2\) implementation for biased docking is freely available and ready to use for any VS campaign as a part of AutoDockTools (http://cbs.scripps.edu/mgltools/ or http://autodockbias.wordpress.com). A complete user guide with a tutorial that thoroughly explains how to apply biases coming from previous knowledge such as ligand-derived pharmacophore or from MD-derived solvent sites is delivered with the program.

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**Notes**

The authors declare no competing financial interest.

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**ABBREVIATIONS**

AmpC, AmpC β-lactamase; AUC, area under the ROC curve; BEDROC, Boltzmann-enhanced discrimination of ROC; BRD4, bromodomain-containing protein 4; CDK2, cyclin-dependent kinase 2; DHFR, dihydropyrimine reductase; DUD-E, directory of useful decoys, enhanced; EF1%, enrichment factor at 1%; EF10%, enrichment factor at 10%; FXa, Factor Xa; FGFr1, fibroblast growth factor receptor 1; Gal-3, galectin-3; HIVpR, human immunodeficiency virus type 1 protease; log AUC, adjusted area under the semilogarithmic ROC curve; MCS, maximum common substructure; MD, molecular dynamics; PDESA, phosphodiesterase 5A; rmsd, root mean square deviation; ROC, receiver operating characteristic; VS, virtual screening


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