ESSENCE-Dock: A Consensus-Based Approach to Enhance Virtual Screening Enrichment in Drug Discovery

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Abstract

Developing new drugs is an expensive and lengthy endeavor, partly due to the reliance on high-throughput screening (HTS), which involves significant costs and is time-consuming. Virtual screening, particularly molecular docking, offers a more cost-effective and faster alternative for identifying promising drug candidates. However, the effectiveness of molecular docking can vary greatly, which has led to the use of consensus docking approaches. These approaches combine results from different docking methods to improve the identification of active compounds and can reduce the occurrence of false positives. However, many of these methods do not fully leverage the latest advancements in docking technology.

In response, we present ESSENCE-Dock (Effective Structural Screening ENrichment ConsEnsus Dock), a new consensus docking workflow aimed at decreasing false positives and increasing the discovery of active compounds. By utilizing a combination of novel docking algorithms, we improve the selection process for potential active compounds. ESSENCE-Dock has been made to be user-friendly, requiring only a few simple commands to perform a complete screening, while also being designed for use in high-performance computing (HPC) environments.

Introduction

The development of novel drugs is a very long and expensive process: depending on the context, the costs of bringing new drugs to market can span from several hundred million to over 4 billion U.S. dollars (1). Many factors contribute to these high costs, including clinical trials and toxicity studies among others. Another part of the high costs can be attributed to hit discovery through expensive, time-consuming high-throughput screening (HTS) (2). Screening small libraries in this manner is done routinely, but chemical space is vast, and navigating it using HTS exclusively can be very costly, both in time and money.

Virtual screening techniques are capable of computationally identifying potential hit compounds, thereby lowering the number of compounds to test experimentally. This can be done much faster and cheaper than HTS, and thus applied to many more compounds, allowing the exploration of a larger chemical space (3). In short, virtual screening techniques are valuable tools for lowering costs and speeding up novel medicine development. Many different types of virtual screening methods exist, but one of the most used techniques is molecular docking (4). For this structure-based screening method, a 3D structure of both the target protein and ligand needs to be available. The goal of molecular docking is to find the optimal ligand pose, which typically has an associated docking score. Subsequently, this docking score can be used to compare different ligands and rank them to obtain a list of the most promising candidates.

The performance of docking algorithms can vary significantly across different targets (5). Examples of popular docking methods include AutoDock Vina (6), and its subsequent forks Smina (7) and Gnina (8), but also commercial software such as LeadFinder (9) and Glide (10) are routinely used in both industry and academia. These methods can be effective on their own, but a consensus docking approach has been shown to improve reliability. By implementing a consensus approach, one can reduce false positives and improve the overall accuracy of results (11). However, many of these existing consensus methods rely on older docking techniques with lower accuracy. Furthermore, they often overlook the importance of the predicted pose similarity, despite its demonstrated effectiveness (12).

To provide a brief overview of existing consensus approaches, we begin with the early yet effective work by A. Kukol (13). In this work, different docking rankings from different algorithms were combined in order to determine which combination of docking methods is the most effective. Specifically, the Directory of Useful Decoys (DUD) (14) was used for this purpose. It was reported that AutoDock 4 (15) combined with AutoDock Vina yielded the best results overall. Houston and Walkinshaw (16) built on these findings by incorporating a simple form of binding pose similarity for additional consensus. Their relatively simple consensus docking method was quite effective at lowering false positives. Their method started by performing molecular docking using both AutoDock Vina and AutoDock 4. Subsequently, they quantified the binding poses of each ligand. In case the RMSD was larger than 2Å, the molecule was rejected and filtered out from further calculations due to the disagreement between the two docking methods. If the RMSD was smaller or equal to 2Å, the compounds were kept for further analysis and ranking. Although simple, this filter step based on RMSD pose-similarity managed to identify many of the decoy compounds, thus demonstrating its value in the whole protocol.

After these developments, consensus docking has continuously been further explored by many researchers. A brief overview of this has been provided in a review of molecular docking written by Torres et al. (11). Some of the most recent consensus docking publications include DockECR (17) and MILCDock (18). Palacio-Rodríguez et al. described an exponential consensus ranking (ECR) method that uses an exponential distribution for each individual rank of every docking method (19). Their method is implemented in such a way that compounds that score very well in a single method can still be ranked highly, even though they score worse in the other docking

methods. The DockECR publication is an in-depth exploration of the ECR method and was collaboratively written with one of the original authors of the exponential consensus ranking paper. They developed an open-source program that has implemented the ECR method and supports several docking methods, such as LeDock (20), rDock (21), Smina (7), and AutoDock Vina (6). Additionally, they added a binding pose similarity metric, which was shown to improve the results in most cases. Their protocol also supports the use of parallelization, enabling simultaneous docking runs across multiple CPUs. Finally, one of the latest works regarding consensus docking is MILCDock. Here, the authors describe a consensus docking method that is enhanced by machine learning. To gather training data, the authors performed molecular docking on datasets such as DUD-E (22) using various open-source docking algorithms including AutoDock Vina, AutoDock 4, and rDock. Subsequently, they used the resulting docking data and binding pose similarities to train machine learning models that can be used as a scoring function. Their results show substantial improvements when compared to the individual docking runs.

Inspired by this prior research, we developed ESSENCE-Dock: a consensus docking workflow primarily focused on reducing false positives and enhancing the enrichment of active compounds. High consensus scores from our protocol correlate with a higher likelihood of compound activity, as demonstrated by our reported high enrichment factors across various DUD-E targets. Our workflow is flexible, user-friendly, and able to be performed using High-Performance Computing (HPC) clusters. The entire screening process is streamlined to just four commands: one for each of the individual docking methods —DiffDock (23), Gnina (8), and LeadFinder (9) — and a fourth to initiate the consensus calculation that integrates the results.

By combining docking scores, binding pose similarity, and ligand flexibility, ESSENCE-Dock generates a comprehensive consensus score to rerank the screened compounds. Additionally, all of the top results can be output to an interactive PyMol Session (PSE) file (24) for visual inspection and user convenience. Furthermore, predicted protein-ligand interactions for the top results are calculated and shown using PLIP (25). These protein-ligand interaction results are also output in tabular and JSON formats for easy integration into further post-docking analysis.

In summary, we introduce ESSENCE-Dock (Effective Structural Screening ENrichment ConsEnsus Dock), a novel consensus docking workflow that addresses some of the identified gaps in current consensus docking methods. ESSENCE-Dock leverages state-of-the-art docking techniques and takes into account binding pose similarity and ligand flexibility to calculate a final consensus score. Our workflow is easy to use, only requiring a few commands for the whole protocol. It is also able to run on HPC clusters using Singularity and has the potential to streamline drug discovery efforts by focusing resources on acquiring and testing compounds with high consensus scores.

Materials and Methods

As established in the introduction, consensus analysis of docking techniques can substantially improve the outcome of a virtual screening campaign (4). With this in mind, we set out to develop our own consensus docking procedure, with a focus on lowering false positives in order to obtain a high enrichment in the top fraction of high-scoring compounds.

As a first step, we attempted to select different types of docking techniques. We selected methods that make use of very different sampling methods for pose generation and use different types of scoring functions. This makes it more significant when consensus, both in high docking scores and binding poses, is found because the docking algorithms came to the same conclusion by employing completely different methods. We ended up selecting LeadFinder (9), Gnina (8) and DiffDock (23). These docking methods, along with their specific sampling and scoring algorithms are discussed in more detail below. A schematic overview of the general workflow is provided in Fig. 1.



Final ESSENCE-Dock results

Fig. 1: Overview of the ESSENCE-Dock Workflow. Each ligand is prepared for and docked using LeadFinder, DiffDock, and Gnina. The resulting docking scores and binding poses are subsequently combined into a final ESSENCE-Dock score, which is used for ranking the compounds.

Ligand and Protein Preparation

Proper preparation of protein and ligands is a crucial step when performing molecular docking, and helps to ensure optimal results. Each of the docking techniques used in our work - LeadFinder (LF), DiffDock (DD), and Gnina (GN)- requires distinct formats for both ligands and proteins. LeadFinder requires the use of mol2 format for both ligands and proteins, DiffDock performs best with ligands in SDF format and protein structures in PDB format, while Gnina

provides optimal results with structures in pdbqt format for both proteins and ligands. This section will detail the procedures used to prepare the ligands and proteins for all the docking calculations, starting with ligand preparation.

Ligands

The selected DUD-E target sets were downloaded from the DUD-E database, with the files 'decoys_final.ism' and 'actives_final.ism' serving as the starting point for conversion (see Fig. 2).



Fig.2: Ligand Preparation Workflow.

Input compounds in SMILES format were processed and converted to 3D SDF, mol2, and pdbqt files for docking with DiffDock, LeadFinder, and Gnina, respectively.

Using Python and RDKit (26), the smiles were processed, hydrogens were added, and a 3D conformation was generated for each compound using RDKit's ETKDG method. The molecules were saved as separate SDF files. Each file was named according to the ID provided in the original .ism files and was tagged as 'active' or 'inactive' for more efficient post-processing after docking.

These SDF files were used as ligands for the DiffDock method but were also converted to mol2 files using ChemAxon's molconvert tool (27). This generated 3D coordinates of a low-energy conformer using the MMFF94 forcefield (28). This made the molecules ready to be docked directly using LeadFinder.

Subsequently, the mol2 files were used as input for the conversion to pdbqt format using the "prepare_ligand4.py" script from AutoDockTools (ADT) (15). Finally, these resulting pdbqt files were used for the Gnina docking procedure.

Protein Structures

For the protein preparation of the DUD-E targets, the PDB code of the provided receptor.pdb file was used to obtain the final receptor files. A schematic overview of the protocol is shown in Fig. 3.



Fig. 3: Protein Structure Preparation Workflow.

Protein structures are imported using their PDB code in Maestro. After processing, the structures are saved as both PDB and mol2 files, which are used for DiffDock and LeadFinder, respectively. Additionally, the PDB file is converted to pdbqt format using AutoDockTools for Gnina docking.

The first step after importing the PDB structure in Schrödinger's Maestro (version 2020-4) (29) was to remove all solvents, ions, and ligands. Subsequently, the Protein Preparation Wizard in Maestro was used to properly (re)assign bond orders, add missing hydrogens, create disulfide bonds, and fill in missing side chains. Next, Maestro's system builder was used to charge the protein using the OPLS3e forcefield (30). Concretely, this was accomplished by minimizing the box volume, defining the solvent model as none, the box shape as orthorhombic, and 10 Å buffer along every dimension.

After these steps, the processed protein was saved in both mol2 and pdb format. The mol2 file was directly used as the receptor file for the LeadFinder docking calculations, while the pdb file was used as input for DiffDock. The pdb file underwent further processing to create a pdbqt file using AutoDockTools (ADT) from MGLTools (version 1.5.7) (15). Concretely, hydrogens were added with the settings "All Hydrogens" and "noBondOrder", followed by assigning AD4 type to all atoms. As a last step, the Gasteiger charge was computed, and the resulting file was saved as a pdbqt file for use in the Gnina docking process.

Docking Techniques

In this section, we will briefly discuss the docking techniques used for our ESSENCE-Dock workflow, detailing their methodology for pose sampling and scoring functions, as well as the parameters we used to process the input ligands.

LeadFinder

LeadFinder (LF), developed by BioMolTech and distributed by Cresset, uses a speed- and accuracy-enhanced genetic algorithm for ligand conformation generation (9). This algorithm creates a diversified pool of conformations, which are subsequently ranked using the LF scoring

function. The output includes the best compounds with their docking score, predicted binding energy, and corresponding docked pose in mol2 format. LF's scoring function is semi-empirical and explicitly accounts for various types of molecular interactions (9).

We ran the LF calculations using MetaScreener v1.1 (31), an open-source and publicly available tool that allows a user to perform molecular docking on high-performance computing (HPC) clusters. This meant that LF could run in parallel, reducing the overall run time substantially and making it feasible to screen larger libraries (see section Run times in Results). The LF docking was executed using version 2104, along with standard variables and a grid size where X, Y, and Z were all set to 30. As user input, only the docking coordinates, as well as the input ligand and protein structure, were included as input variables. The LF docking calculations were performed on the SCBI UMA Picasso HPC cluster using 1 CPU core and 2 GB of RAM per job.

Gnina

Another docking method we used in our consensus docking approach is Gnina (8). Gnina is a fork of Smina (7), which in turn is a fork of Autodock Vina (Vina) (6). In contrast with LeadFinder, which utilizes a genetic algorithm to explore a ligand's conformational space, Gnina employs Monte Carlo sampling to generate a diverse ensemble of ligand conformations. It also implemented an ensemble of convolutional neural networks (CNN) as a scoring function, which outperforms the scoring functions of its predecessors Vina and Smina in most cases (8). Although Vina's scoring function is still used during pose generation and refinement for computational efficiency, Gnina's CNN ensemble model rescores and sorts the final poses. These settings, found to be the most efficient by the authors, were also used for our Gnina docking calculations.

For our calculations, the Gnina docking algorithm was implemented in MetaScreener v1.1, again providing support to run the calculations massively in parallel on HPC clusters. Because of the CNN scoring function, Gnina is able to run a lot faster on a GPU compared to a CPU. Despite this, the parallel execution of calculations on an HPC cluster made processing the selected DUD-E datasets still feasible. Gnina Version 1.0 was used with standard settings, a box size of 30 in all dimensions, along with the user-provided ligand, protein, and docking coordinates. Like the LF calculations, the GN docking calculations were performed on the SCBI UMA Picasso HPC cluster, but using 4 CPU cores and 2 GB of RAM per job.

DiffDock

The final docking technique we integrated into our consensus docking workflow is DiffDock (23). DiffDock is a novel, state-of-the-art blind docking method based on geometric deep learning. The process refines randomized ligand conformations at various protein locations via a trained reverse diffusion process, simultaneously determining the best binding pocket and conformation. This eliminates the need for a pre-designated search box as required by LF and Gnina. Because this is a blind docking method, the whole protein and all its potential binding pockets are explored.

Unlike the other methods, DiffDock does not provide a scoring function. It does offer a confidence score indicating prediction certainty. Although this is useful, it still does not allow a user to gauge how potent a ligand binds, making it difficult to rank different ligands accurately

among each other. Still, DiffDock is capable of predicting a ligand's binding pose with state-of-the-art accuracy, so it is still very useful to compare it to the binding poses generated by LeadFinder and Gnina.

To facilitate DiffDock use in virtual screening, we developed a fork of DiffDock named DiffDockHPC. It enables multiple parallel DiffDock jobs, significantly speeding up calculations when processing more extensive libraries, and freely available at https://github.com/Jnelen/DiffDockHPC. Like Gnina, DiffDock can be greatly accelerated by GPUs, but running the calculations massively in parallel on CPUs still makes it feasible to be a part of the consensus docking workflow. The DiffDock calculations were performed on NLHPC's leftraru cluster, using 4 CPU cores and 8 GB of RAM per job.

Processing of Docking Results

In this section, we describe the processing of docking results, the details of our scoring formula, and the final output files that our ESSENCE-Dock workflow generates.

Scoring Formula

To easily rank compounds, we developed a scoring formula that considers not only individual docking scores but also the binding pose similarity, as calculated by the average RMSD between the docking poses. The ligand's flexibility is also taken into account by incorporating the number of rotatable bonds into the equation. The inclusion of the number of rotatable bonds ensures that results are not skewed towards rigid structures, which inherently have fewer degrees of freedom and consequently a higher chance of achieving better binding pose similarity. The ESSENCE-Dock rescoring formula is presented in Equation 1:

$$ESSENCEDockScore = \sqrt{\frac{2}{RMSDaverage}} \cdot MBE \cdot \sqrt{ln(RB + 1) + 1}$$
(1)

Here, RMSDaverage denotes the average root mean square deviation (as shown in Eq. 2), and MBE represents the Mean Binding Energy (computed as per Eq. 3). RB corresponds to the number of rotatable bonds. The RMSD and the number of rotatable bonds are calculated using OpenBabel's obrms and obrotamer functionalities respectively (32).

$$RMSDaverage = \frac{RMSD_{DD_{GN}} + RMSD_{DD_{LF}} + RMSD_{GN_{LF}}}{3} (2)$$
$$MBE = \frac{BindingEnergy_{GN} + BindingEnergy_{LF}}{2} (3)$$

As can be derived from Eq. 1, the ESSENCE-Dock score is inversely proportional to the agreement between the binding poses produced by the individual docking methods. This means that a lower (more negative) ESSENCE-Dock score indicates a higher degree of agreement between the docking methods, and therefore a more reliable result.

Note as well that in equation 1, just the average RMSD and docking scores and the number of rotatable bonds are required. In this sense, our approach is flexible and able to be used with other docking programs as well. For user convenience, the whole ESSENCE-Dock approach

has been implemented in Metascreener v1.1, and the consensus rescoring and post-processing can be executed using just a single command

Enrichment Factors

To validate the protocol, we selected a diverse set of DUD-E targets and computed enrichment factors (EFs) at different thresholds. We compared these EFs with those obtained from individual docking methods (LeadFinder and Gnina) as well as a more basic consensus docking approach using the mean binding energy and mean rank as scoring metrics. The EF for the top 5, 1, 0.5, and 0.1% was calculated using Eq. 4:

 $EF = \frac{activesEF\%/totalEF\%}{AllActives/AllCompounds}$ (4)

At its core, this formula compares the fraction of actives in the top x% of compounds to the overall distribution of actives in the dataset. ActivesEF% represents the amount of actives in that EF% fraction, while total EF% represents the total number of compounds part of the same EF% fraction. The bottom part of the equation is the fraction of the total amount of actives and the total amount of compounds of the whole dataset.

Output Files

After scoring and ranking all the compounds using Eq. 1, the workflow is able to generate an interactive PyMol (24) session (PSE) file. This PSE file contains the protein structure and the top 50 results (user-configurable) based on the scoring.

Each entry in the PSE file displays the three docked structures from LeadFinder, DiffDock, and Gnina, enabling easy visual assessment of binding pose similarity. It also provides the original docking score for LeadFinder and Gnina structures. Additionally, the file includes predictions of protein-ligand interactions as predicted by PLIP (25), offering insights into the potential binding interactions.

The interactive PSE file facilitates the study and comparison of ligand-protein interactions across different docking methods. Furthermore, the PLIP results are summarised in both JSON and spreadsheet-style CSV files for convenient further analysis.

Finally, the complete ESSENCE-Dock information (the individual docking scores, RMSD values, etc.) for all compounds is saved to an output CSV file as well.

Results and Discussion

To demonstrate the efficacy of ESSENCE-Dock, we applied our workflow to a set of 21 distinct DUD-E targets, representing a diverse range of protein families and biological contexts. After docking and ranking the compounds, the enrichment factors (EF) were calculated at different fractions. EFs are a measure of the method's ability to identify active compounds among the top-ranked candidates, with higher EF values indicating better performance (refer to Materials and Methods Eq. 4 for the exact EF formula). We evaluated EFs for the individual DUD-E targets at various fractions: the top 5%, top 1%, top 0.5%, and top 0.1% to analyze

ESSENCE-Dock's efficacy in different scenarios. To compare and benchmark ESSENCE-Dock's performance, we also computed the EFs at identical threshold levels using the rankings from the individual docking methods: LeadFinder (LF) and Gnina (GN). Additionally, we calculated two basic consensus metrics using the individual GN and LF docking calculations: the Mean Binding Energy (MBE), which provides insights into the collective binding affinities, and the Mean Rank (MR), which offers a measure of the ranking consistency across different docking methods. All of these results are summarized in Table 1. The average results are compared using a barchart in Fig. 4. The same information for the Median and all of the results for the individual DUD-E targets can be found in the Supplementary Data (Fig. S1).



Fig. 4: Bar Chart of Average Enrichment Factors (EF) Across 21 Various DUD-E Targets at Varying EF Thresholds.

This figure provides a comparative analysis of Enrichment Factors (EF) across 21 distinct DUD-E targets. The Y-axis represents the enrichment factor, while the X-axis indicates the corresponding EF threshold values. The color-coded bars correspond to the different ranking methods: LeadFinder, Gnina, Mean Rank, Mean Binding Energy and ESSENCE-Dock.

Table 1: Comparative Performance of ESSENCE-Dock and Reference Docking Methods Across 21 DUD-E Targets at Various Enrichment Factors (EF%). Table 1 compares the performance of ESSENCE-Dock (Cons.) with reference methods (LeadFinder - LF, Gnina - GN, Mean Rank - MR, and Mean Binding Energy - MBE) across 21 DUD-E targets at various Enrichment Factors (EF%). The table presents EF values for top 5%, top 1%, top 0.5%, and top 0.1%. Shaded cells indicate method rankings, with green denoting higher comparative EF and red indicating lower comparative EF.

	EF5%					EF1%					EF0.5%					EF0.1%				
	LF	GN	MR	MBE	Cons.	LF	GN	MR	MBE	Cons.	LF	GN	MR	MBE	Cons.	LF	GN	MR	MBE	Cons.
ADA	3.9	3.2	4.3	4.1	5.6	1.1	0.0	9.8	5.4	11.9	0.0	0.0	8.5	4.3	21.3	0.0	0.0	9.9	0.0	39.7
AKT1	1.7	3.6	2.7	3.3	1.4	1.7	8.6	4.4	6.5	2.7	2.7	12.2	6.8	8.8	2.7	0.0	26.9	6.7	16.8	3.4
AMPC	0.8	0.8	0.8	0.0	2.9	0.0	0.0	0.0	0.0	2.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
CASP3	2.9	4.2	5.5	5.0	5.0	9.5	4.5	10.6	11.1	9.5	11.2	4.1	13.2	15.2	17.2	5.0	0.0	14.9	24.9	39.8
CP3A4	2.8	1.5	2.4	2.9	2.0	4.7	1.8	2.3	4.1	1.8	8.2	2.3	4.7	7.0	0.0	11.7	5.9	5.9	5.9	0.0
CXCR4	2.0	0.5	1.0	0.5	0.5	0.0	0.0	2.5	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
FA7	5.6	8.6	9.0	9.3	7.5	8.7	14.8	20.1	20.9	17.4	7.0	17.4	29.7	24.4	22.7	0.0	0.0	55.8	37.2	55.8
FABP4	0.4	8.1	3.4	5.5	6.4	0.0	21.3	2.1	8.5	10.6	0.0	25.5	0.0	8.5	8.5	0.0	0.0	0.0	0.0	0.0
FPPS	0.0	5.6	0.0	0.0	9.4	0.0	2.4	0.0	0.0	20.1	0.0	0.0	0.0	0.0	28.0	0.0	0.0	0.0	0.0	11.7
GCR	0.7	5.3	2.9	3.9	4.7	0.0	15.8	6.2	13.5	13.9	0.0	21.8	7.0	21.0	21.0	0.0	47.3	7.9	23.7	27.6
GLCM	5.5	1.1	3.7	4.1	5.5	11.0	1.8	3.7	11.0	5.5	11.3	0.0	3.8	15.0	7.5	0.0	0.0	0.0	0.0	0.0
HIVPR	5.1	4.5	7.1	7.0	6.5	5.4	6.2	14.2	11.2	9.7	4.5	6.0	17.6	14.2	12.0	3.8	9.4	26.3	24.4	26.3
HIVRT	2.5	3.3	3.8	3.8	4.2	2.7	3.6	5.6	5.6	8.9	1.2	4.7	7.1	5.9	13.0	0.0	9.0	9.0	6.0	18.0
HS90A	1.8	0.0	0.2	0.5	3.9	2.3	0.0	0.0	0.0	8.0	2.2	0.0	0.0	0.0	13.5	0.0	0.0	0.0	0.0	11.2
ITAL	3.3	3.2	2.9	3.3	3.0	8.0	11.6	11.6	13.1	9.5	13.1	8.7	21.8	20.4	17.5	7.0	7.0	55.6	48.7	48.7
KIF11	1.2	8.3	6.9	7.8	7.4	0.0	21.4	8.6	16.3	24.0	0.0	30.9	6.9	17.2	34.3	0.0	25.7	0.0	0.0	25.7
MK14	2.5	3.3	3.3	3.9	2.4	5.2	9.2	6.6	8.3	5.4	5.9	13.5	9.7	12.8	9.3	1.8	31.5	21.0	29.8	29.8
NRAM	0.6	1.2	3.1	2.0	7.5	0.0	0.0	2.0	0.0	17.3	0.0	0.0	0.0	0.0	29.0	0.0	0.0	0.0	0.0	53.6
PA2GA	2.4	5.3	1.6	1.8	8.9	0.0	6.1	1.0	1.0	25.5	0.0	6.1	2.0	2.0	30.6	0.0	10.6	10.6	10.6	31.8
TRY1	3.0	5.9	6.0	6.1	4.0	3.6	6.5	6.7	8.2	8.2	3.1	7.6	7.6	9.4	12.0	6.8	18.1	9.1	9.1	27.2
WEE1	14.3	17.6	15.3	16.8	17.6	41.8	60.3	61.3	61.3	53.5	47.5	61.3	61.3	61.3	59.3	61.3	61.3	61.3	61.3	61.3
Average	3.0	4.5	4.1	4.4	5.5	5.0	9.3	8.5	9.8	12.6	5.6	10.6	9.9	11.8	17.1	4.6	12.0	14.0	14.2	24.4
Median	2.5	3.6	3.3	3.9	5.0	2.3	6.1	5.6	8.2	9.5	2.2	6.0	6.9	8.8	13.5	0.0	5.9	7.9	6.0	26.3

As illustrated in Table 1, our ESSENCE-Dock method often outperformed the other methods, especially at the smaller EF fractions of EF0.5% and EF0.1%. In the case of EF0.1%, ESSENCE-Dock was only surpassed by other methods in five cases: AKT1, CP3A4, GCR, ITAL, and MK14. Even when our approach performed worse, it often still showed notable enrichment. Only in the cases of AKT1 and CP3A4 did ESSENCE-Dock substantially underperform other methods. For the other three DUD-E targets, ESSENCE-Dock's performance remained competitive with high EFs of 27.6, 48.7, and 29.8 for GCR, ITAL, and MK14, respectively. In some other cases such as AMPC, CXCR4, FABP4, and GLCM, none of the methods manage to identify actives in the top 0.5% and 0.1% of ranked compounds, resulting in EFs of 0. However, an important note is that these specific datasets are relatively small (ranging from only ~2900 to ~3850 compounds). This means that at these top enrichments, only a few compounds are considered, which can partly explain why all of the methods score substantially worse here.

The individual EFs were averaged for each method and EF threshold, as summarized in Fig. 4. This figure clearly demonstrates that our approach achieved the highest average enrichment scores across all evaluated thresholds. While the improvement at EF5% is relatively modest, the improvement becomes more evident as the EF threshold decreases. This is especially the case with EF0.1%, where ESSENCE-Dock has an enrichment factor that is 5 times higher than LeadFinder and 2 times higher than Gnina. Furthermore, even in comparison to the Mean Rank and Mean Binding Energy consensus methods, ESSENCE-Dock is still nearly twice as effective.

Fig. 5 shows graphs comparing the ranking performance of the individual docking methods (GN and LF) and the ESSENCE-Dock approach for three different DUD-E datasets: NRAM (A), CASP3 (B), and CP3A4 (C). Each figure within this comparison features two separate histograms: one representing the distribution of active compounds (in blue) and the other depicting the distribution of decoy compounds (in orange). Of note are the varying quantities of active and decoy compounds which lead to separate Y-axes for each group. This distinction is caused by the significantly larger number of decoys compared to active compounds. For all docking methods, lower (more negative) scores on the X-axis indicate a better result. Similar figures for all of the tested DUD-E datasets are provided in the Supplementary Data (Fig. S2).





This figure presents the distributions of actives (depicted in blue) and decoys (represented in orange) based on their respective docking scores for the NRAM (A), CASP3 (B), and CP3A4 (C) datasets included in the DUD-E database. Notably, the figure employs differently scaled axes to accommodate the varying quantity of actives (left axis) compared to decoys (right axis). In all cases, a lower score (more negative) indicates a better-ranked sample.

In the example of NRAM (Fig. 5A), the individual docking methods GN and LF don't excel at assigning better docking scores to the active compounds compared to the decoys. This is reflected in the overlapping distributions between the actives and decoys in these cases. In contrast, the assigned ESSENCE-Dock scores produce a distinct cluster of active compounds distributed within the range of -25 to around -15, clearly separated from the decoy compounds. These trends align with the quantitative data presented in Table 1, where our ESSENCE-Dock

method at EF0.5% and EF0.1% for NRAM demonstrates substantial enrichment, while the other methods yield enrichment factors of zero.

In the case of CASP3 (Fig. 5B), the individual docking methods GN and LF demonstrate better performance compared to the NRAM case, successfully ranking a higher number of actives near the top based on their respective docking scores. Still, ESSENCE-Dock is able to outperform the other methods using the consensus approach. Again, this is reflected in Table 1, where at EF0.5% and especially EF0.1%, ESSENCE-Dock shows a higher EF than the individual docking methods, as well as the simplistic consensus MR and MBE results. Interestingly, again there is a distinct cluster of active compounds present between the -20 and -15 scores.

Finally, Fig. 5C shows the example of CP3A4 from the DUD-E database. This is one of the cases where ESSENCE-Dock performs worse compared to the other methods, as can also be seen in the EF0.1% results for CP3A4 in Table 1. Even the individual docking methods, GN and LF provide better enrichment than ESSENCE-Dock in this specific example. While these results for ESSENCE-Dock on the CP3A4 dataset seem disappointing, they are also informative. The observation that ESSENCE-Dock assigns lower scores to nearly all compounds in this dataset indicates its effectiveness in detecting a lack of consensus, which, in turn, suggests that the docking results may be unreliable.

In this context, the ESSENCE-Dock score can also be used as a confidence metric for the consensus results. When the score is relatively bad (for example >-12.5), it usually means that ESSENCE-Dock detects substantial disagreement between the individual docking methods. This suggests that the consensus results are relatively unreliable and that caution should be taken when interpreting them. This is supported by the results for most other datasets where ESSENCE-Dock performs poorly (AMPC, CP3A4, CXCR4, FABP4, and GLCM), which typically have low ESSENCE-Dock scores (>-15), indicating low confidence. We believe that the ability of ESSENCE-Dock scores being able to act a quality measure for the docking results is a valuable feature. Concretely, during virtual screening campaigns using ESSENCE-Dock, a certain score cutoff (for example -15 or -20) could be chosen. Compounds scoring worse than the threshold would be rejected, allowing researchers to effectively sift through and prioritize compounds, reducing false positives and focusing on candidates with a more robust consensus.

Finally, we briefly discuss the runtimes of three DUD-E examples: MK14, TRY1 and WEE1. These datasets vary in size, and can thus give an indication for the runtimes across different dataset sizes. We will discuss the runtimes of the individual docking runs (Fig. 6), as well as the time it takes to process the individual docking results into the final ESSENCE-Dock rescoring (Fig. 7).



Fig. 6: Wall-Clock Runtime Performance of Docking Algorithms on DUD-E Targets. Bar chart depicting wall-clock runtimes for DiffDock, Gnina, and LeadFinder applied to MK14, TRY1, and WEE1 datasets, highlighting the total wall-clock runtime of each method across varying dataset sizes.

Fig. 6 summarizes the total runtimes for the individual docking methods: DiffDock, Gnina, and LeadFinder, calculated for the three aforementioned DUD-E datasets. MK14 included 36,428 compounds, TRY1 had 26,429, while WEE1 contained 6,252. DiffDock calculations were performed on the NLHPC Leftraru supercomputer ("slims" node) with 4 CPU cores and 8GB RAM per job. In contrast, Gnina and LeadFinder docking calculations were carried out on the SCBI UMA Picasso HPC cluster, with Gnina being assigned 4 CPU cores and 2GB RAM, while LeadFinder using 1 CPU core along with 2GB RAM per job.

In all cases, the Gnina calculations took the most amount of time. This can probably be attributed to the fact that these calculations were run using only CPUs, while Gnina's CNN scoring algorithm has been optimized to be run with a GPU. The Gnina docking calculations were assigned 4 CPUs per job, which was able to accelerate the initial generation and optimization of binding poses compared to running it with only 1 CPU. However, the final step of CNN scoring took up the most amount of time, and seemed to be unchanged regardless of the amount of assigned CPUs. Thus, it is probably not able to take advantage of multiple CPU cores during the CNN scoring calculations. However, access to GPUs could accelerate the CNN scoring and reduce overall runtime substantially.

In most cases, DiffDock and LeadFinder took about the same amount of wall-clock time to complete. The exception here is with TRY1: here the LeadFinder calculations took even longer than in the case of MK14. We assume this is an anomaly, as this is not in line with the trend that we observed with the other datasets. One potential explanation could be due to a high server load during those specific calculations, leading to overall longer runtimes.

Even though their runtime was similar, DiffDock and LeadFinder don't have the same computational cost: DiffDock was assigned 4 CPU cores, while LeadFinder ran on only one. Note that DiffDock was also designed to be run on a GPU, but when it is not, it is still able to take advantage of multiple CPU cores if they are available (unlike Gnina). Finally, LeadFinder is made to be run on CPUs and thus is the most optimized. This translated into the lowest runtimes in all scenarios, except in the anomalous case of TRY1. Note that the total runtime is quite large, but when distributing this total time across many jobs (100 to even 250 jobs) on HPC clusters, the total wall-clock runtime still is reasonable. However, screening very large libraries (1 million compounds or more) would be a big task. Hence we recommend to first perform a prescreening, using for example faster ligand-based techniques to narrow down the amount of compounds. For this purpose, one could employ techniques such as fingerprint similarity, shape-based screening or pharmacophoric similarity screening among others.





Fig. 7: Runtime comparison of the ESSENCE-Dock Consensus calculations across DUD-E targets with varying sizes.

Scatterplot and trendlines depict UNIX real, user, and system runtimes for ESSENCE-Dock across WEE1, TRY1, and MK14 datasets, with R² values indicating highly linear nature when increasing the dataset size.

In Fig. 7 we show the runtimes of the ESSENCE-Dock consensus calculations. In this process, the individual docking calculations are processed and combined into the final ESSENCE-Dock. The time to generate a PyMol Session file with PLIP scores is not included here. These consensus calculations are performed using a Python script, which also makes use of subprocess calls to obrms and obrotamer to calculate the RMSD and retrieve the number of rotatable bonds. The time spent in subprocess calls is not included in the user time. Instead it is listed as system time and therfore we include all 3 metrics as returned from the Unix "time" command. These calculations were performed on a local workstation using 1 core and 2GB of RAM on an AMD EPYC 7282 16-core processor.

Fig. 7 also shows that the ESSENCE-Dock consensus calculations scale linearly. The trendlines have an R² value of 0.99 for system-time and 1.00 for user- and real-time, which confirms its linear relationship. The total runtime can get more significant if the datasets are really large, but it is still manageable, only taking up about 15 minutes for a small dataset (like the WEE1 example) and about an hour and 10 minutes for larger ones (like the MK14 example).

From these results, it is clear that our novel ESSENCE-Dock approach is a powerful docking technique that is able to provide substantial enrichment across various families of proteins. On top of that, it also provides helpful output files, including a PSE file of the top results along with their predicted PLIP interactions. The predicted protein-ligand interactions are also summarized in a useful JSON and CSV file.

Conclusions and Outlook

We have presented ESSENCE-Dock, an easy-to-use consensus docking workflow that focuses on providing a high enrichment of active compounds. It combines three different docking methods (DiffDock, Gnina, and LeadFinder) and considers the predicted binding pose similarities, docking scores, and ligand flexibility. Additionally, the top results can be visualized in an interactive PyMol session with PLIP-predicted protein-ligand interactions. The predicted protein-ligand interactions are also provided in useful formats for post-processing. ESSENCE-Dock can be run on HPC clusters using Singularity and Slurm, making it feasible to apply the workflow on larger virtual libraries.

Future work could explore the use of new docking methods, as ESSENCE-Dock is flexible and can easily be made to work with other docking methods. When new, better-performing docking methods are developed, they can be quickly and easily integrated into the workflow. Additionally, it could be interesting to train a machine-learning algorithm on all of this input data. This would allow us to move to a machine learning scoring function, which could be more robust and outperform the current method, leading to even better overall results.

Data and Software Availability

DiffDockHPC: <u>https://github.com/Jnelen/DiffDockHPC</u> Metascreener: <u>https://github.com/bio-hpc/metascreener</u> The DUD-E docking results and consensus: <u>https://zenodo.org/doi/10.5281/zenodo.10025839</u>

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Abbreviations

ADT: AutoDock Tools CNN: Convolutional Neural Network Cons.: Consensus CPU: Central Processing Unit CSV: Comma Separated Values (file) DD: DiffDock DUD(-E): Directory of Useful Decoys(-Enhanced) ECR: Exponential Consensus Ranking **EF: Enrichment Factor** GN: Gnina **GPU:** Graphics Processing Unit HTS: High-throughput Screening HPC: High-Performance Computing LF: LeadFinder MBE: Mean Binding Energy ML: Machine Learning MR: Mean Rank PDB: Protein Data Bank PLIP: Protein-Ligand Interaction Profiler PSE: PyMol Session (file) RAM: Random Access Memory **RB:** Rotatable Bonds **RMSD: Root Mean Square Deviation VS: Virtual Screening**