# Cosolvent and Dynamic Effects in Binding Pocket Search by Docking Simulations

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### Introduction

Proteins are ubiquitous building blocks playing a critical role in the reproduction, metabolism, and regulation of living organisms and viruses. Understanding and manipulating the way proteins interact with their surrounding is, therefore, of utmost interest from both a biological and a medical point of view. Currently, the most important method to manipulate the function of proteins is through the administering of drugs. For this reason, there exists a growing interest in identifying new binders for a wide variety of proteins in the hopes of treating a number of different sicknesses. <sup>1–7</sup> In fact, 78 % of the biological drugs approved by the United States Food and Drug Administration (FDA) have clear protein molecular targets. <sup>8</sup> Therefore, it is not surprising that scientists turned to them once again, when faced with the new and immediate challenges of the coronavirus disease 2019 (COVID-19)

pandemic.

The COVID-19 pandemic caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) continues to claim thousands of lives every day more than a year after its outbreak. However, the knowledge about it and the developed tools to fight against it are vastly more potent than they were a year before. 10 Antiviral drugs targeting the proteins vital to the reproduction of SARS-CoV-2 have been the most important tools, aside from vaccines which can only be used as preventative measures. For example, Remdesivir, one of the most widely used antiviral drugs against SARS-CoV-2 around the world, 11 targets the RNA-dependent RNA polymerase (RdRp) protein of the virus. 12 Furthermore, given the urgency of developing an effective treatment, most attempts to find new inhibitor substances were in fact drug repurposing studies, targeting the virus's RdRp<sup>13-17</sup> or other important proteins. <sup>18–21</sup> The RdRp protein is an especially promising drug target as it is responsible for the replication of the viral RNA inside the host cell, 22 and it is highly similar to the RdRp of SARS-CoV, 23 which already has a number of verified inhibitors. 24 In addition, its high-quality three dimensional (3D) structure has been available from as early as April 2020.<sup>25</sup> In large part due to the urgent nature of the COVID-19 pandemic, most of the above cited research projects relied heavily, or even exclusively, on computational techniques for the discovery of the potential inhibitors, due to the cost and time efficiency of such methods.

High-throughput screening enables the routinely evaluation of thousands of substances in a week. <sup>26</sup> This tremendous efficacy is often supported by the development and application of innovative computational methods, which became more useful since the advent of structure-based drug design, where potential drugs are created or found based on the 3D structure of the protein target. <sup>27,28</sup> Although such target structures were initially only obtainable through costly and cumbersome experimental methods, such as X-ray crystallography <sup>29</sup> or nuclear magnetic resonance (NMR) spectroscopy, <sup>30</sup> they are nowadays much more readily available due to the gradual improvement of existing methods, the appearance of new experimental methods, such as cryo-electron microscopy, <sup>31</sup> and the development of recent computational

techniques, such as homology modeling.<sup>32</sup> Taking advantage of the quickly growing body of available genomic data, computational tools capable of predicting protein structures from mere amino acid sequence information have also been developed.<sup>33,34</sup> By employing one (or a combination) of the above techniques, high-quality structures are available for a larger number of protein targets than ever before.

The current challenge to computational chemists is therefore how to best utilise the available structural information. The computational methods developed for structure-based drug design fall into two main categories: de novo design methods construct new, tailored ligands, while docking methods select ligands complimentary to the target from the existing compound space. Among the docking methods, virtual screening (VS) has emerged as a particularly successful technique. This procedure can be thought of as a computational extension to high-throughput screening, where a large number of compounds are docked to the target protein structure in silico. Traditionally, VS campaigns have been carried out utilising a single, experimentally determined protein structure, often in the crystallised form. The deficiencies of using only a single crystallised protein structure has been recently recognised. Tirstly, the structure of the crystallised protein often differs significantly from the conformations that the protein adopts in vivo. Secondly, even if the crystal structure is representative of the conformation most often visited in solution, a single structure cannot account for the dynamics of protein motion.

Different theoretical models that consider the importance of protein motion have been developed, e.g., the induced-fit model of ligand docking,  $^{41,42}$  where the structure of the protein may change during ligand uptake, or the model of conformational selection,  $^{43-45}$  which views the target protein as a dynamic object even in the absence of ligands. The need to take protein flexibility and motion into account became even clearer with the discovery of cryptic or hidden pocket structures.  $^{46-48}$  The characteristic property of these pockets is that they only appear in the presence of the appropriate ligand, while their existence is not obvious from the equilibrium structure of the protein. The exact mechanism of their formation is not

yet clear, although some combination of induced-fit and conformational selection has been hypothesised.<sup>47</sup> The discovery and theoretical description of such pockets are hindered by the fact that their opening often requires large scale rearrangements of the protein structure, events that are traditionally hard to predict with computational techniques.<sup>49</sup>

With the importance of protein dynamics gaining wider recognition, new, more elaborate methods are appearing which aim to account for this phenomenon. On the one hand some of the modern computational docking programs, such as AutoDock Vina, <sup>50</sup> can treat a selected number of protein residues as flexible at the cost of increased calculation times. This method is well suited to study a previously known, specific binding site of the protein. However it cannot account for larger structural changes of the protein and is limited to a handful of flexible residues due to its computational requirements. On the other hand, the family of ensemble docking techniques utilises traditional (rigid protein) docking calculations in combination with an ensemble of protein conformations to account for the flexibility of the target. <sup>35,37</sup> The careful selection of the structures of the ensemble can enable the description of large scale conformational changes and to the discovery of new cryptic pockets. <sup>45,48,49</sup> The main challenge for these methods is the generation of the protein structure ensemble, which can be achieved experimentally by using different crystallised structures <sup>35,51,52</sup> or computationally by, e.g., conformational space searches, <sup>53</sup> neural networks <sup>54</sup> and molecular dynamics (MD). <sup>37,55</sup>

MD is an especially promising avenue, after all it has been designed for the very purpose of efficiently sampling the realistic conformational space of proteins. However, one of the largest obstacle of MD calculations is the extremely slow convergence of the calculated trajectories, <sup>37</sup> which precludes the population of rarely visited conformations. Even with highly specialised code and computers the longest timescales reachable are in the range of milliseconds. <sup>56</sup> In order to be able to sample rare events, a number of modified MD techniques have been developed. The first group of these is the enhanced sampling methods, where some unphysical bias is introduced into the simulation in order to encourage the

sampling of otherwise unlikely conformations. Some of the most popular enhanced-sampling methods in the context of cryptic pocket discovery are umbrella sampling, <sup>57</sup> steered MD, <sup>58</sup> metadynamics, <sup>59</sup> and replica exchange MD, <sup>60</sup> among others. A completely separate approach for the sampling of rarely visited conformations harboring cryptic pockets is that of the cosolvent methods. The main idea behind these frameworks is to replace the traditional water solvent in MD simulations with a mixture of water and some other cosolvent. The oftentimes hydrophobic or amphipatic cosolvent probes can then interact with the protein and occasionally induce conformational changes or stabilise some conformations where a cryptic pocket is open. Cosolvent methods have been successfully used to identify cryptic sites in a number of targets. <sup>45,48,61,62</sup>

The primary aim of the presented work is investigate the effect of protein dynamics in the results of a VS campaign. The ensemble of protein structures is obtained via MD simulations. Further sampling is obtained by cosolvent trajectories where water/benzene and water/phenol mixtures are employed. Recognising the severity of the COVID-19 pandemic, the calculations are carried out on the RdRp protein of SARS-CoV-2 and a set of FDA approved small molecule drugs, in the hopes of contributing to the generation of knowledge necessary to develop effective treatments against this virus.

## Computational Details

The SARS-CoV-2 RdRp protein complex was chosen as the target of our investigations. In its active form it is composed of three domains: nonstructural proteins 7, 8 and 12 of SARS-CoV-2. Its active site is located in a deep groove and is highly similar to that of the analogous protein of the SARS-CoV.<sup>24</sup> Its simulation ready structure was obtained from the website of D. E. Shaw Research,<sup>63</sup> where extensive MD simulations have already been carried out for it. In Reference 13, the authors note that two zinc ions are necessary for the structural integrity of the protein. These ions were however not found in the structures

and trajectories downloaded from D. E. Shaw Research. After numerous failed attempts at stabilising these zinc ions in their bound positions with restraining potentials and gradual heating, their inclusion was rejected in favor of the original D. E. Shaw structure. Additionally, two crystal structures determined with cryo-electron microscopy were downloaded from the Protein Data Bank website: the apo structure 6M71<sup>64</sup> and the holo structure 7B3B.<sup>65</sup> Out of these two crystallised structures, only the apo structure was utilised for docking calculations, to emulate drug discovery VS campaigns where the holo structure of the target is not available. The holo structure was used only to visualise the conformational changes around the active site occurring during ligand binding as will be explained below.

The MD calculations were carried out with the Amber 18 program package, <sup>66</sup> according to the following protocol. Three types of solvent boxes were prepared for the simulations: a simple water one and two with either benzene or phenol as cosolvent. The protein structures were solvated in octahedral solvent boxes containing the appropriate mixture of water and cosolvent molecules. A distance of at least 12 Å was left between the protein and all sides of the solvent box. The charge of the system was neutralised with sodium ions. During the simulations, periodic boundary conditions and a 12 Å cutoff for the Lennard–Jones interactions were used. The solvated systems were first minimised for 1000 gradient descent steps followed by an other 1000 conjugate gradient steps. Next, the heating of the systems to 300 K were performed during a 1 ns simulation with the Langevin thermostat in the NVT ensemble. Finally, 200 ns production simulations were carried out at 300 K and 1 bar pressure using the Langevin thermostat and Berendsen barostat in the NPT ensemble. Three replicas were run for the production calculation for each solvent. The last two simulations for each solvent were started from a random equilibrated frame of the first simulation for that solvent, with the velocities of all particles randomised according to the Boltzmanndistribution. The production calculations were run with GPU acceleration, using the pmemd program of Amber 18.

During the preparation of the solvent boxes containing cosolvents the packmol program

was utilised.  $^{67}$  The concentration of the cosolvents were set to 10 v/v % in both cases. In the case of the benzene cosolvent severe clustering of the cosolvent molecules was observed during the MD simulations when the default force field parameters were used. To circumvent this issue, scripts included in the ParmEd distribution  $^{68}$  were utilised to introduce Lennard–Jones potentials between the carbon atoms of different benzene molecules.

The exact form of this artificial potential between carbon atoms i and j is:

$$V_{ij} = \gamma \left[ \left( \frac{R_{\min}}{r_{ij}} \right)^{12} - 2 \left( \frac{R_{\min}}{r_{ij}} \right)^{6} \right]. \tag{1}$$

Here,  $V_{ij}$  is the introduced LJ potential,  $\gamma$  is the parameter determining the minimum value of the potential,  $R_{\min}$  is the parameter controlling the position of the minimum, while  $r_{ij}$  is the distance between carbon atoms i and j. The default parameter values of  $\gamma = 0.00036$  kcal/mol and  $R_{\min} = 7.12719$  Å were utilised. After this modification was made, no clustering of the benzene molecules was observed during the simulations.

#### Trajectory clustering

For the clustering of the MD trajectories the cpptraj program <sup>69</sup> of Amber 18 was utilised. A density based clustering algorithm (chosen with the dbscan keyword of cpptraj) was employed, with the parameters k (unitless) and  $\varepsilon$  (in ångströms) set to 4 and 1.1 Å respectively (see Section ?? for the discussion of this choice). For each type of solvent the equilibrated part of the trajectories of the three replica simulations were concatenated and the clustering was carried out separately for each solvent. Before clustering, the structures in every frame were aligned to each other by their alpha carbon atoms. The clustering was performed using the RMSD values of the alpha carbon atoms as the distance metric between the conformations. A total of 19 cluster representatives were obtained with 13 coming from the trajectory with water as the solvent while the benzene and phenol cosolvent trajectories yielded 3 cluster representatives each.

#### **Docking calculations**

The set of FDA approved drugs were downloaded from the ZINC database <sup>70</sup> in the mol2 format. This set is a popular choice for drug repurposing studies <sup>15–17</sup> and with approximately 2000 thousand contained ligands, it was feasible to perform docking calculations for all protein conformation, ligand pairs. From this set, 1957 ligand structures were converted to the pdb format, necessary for docking with AutoDock Vina, with the openbabel program. <sup>71</sup> The 19 cluster representative protein structures along with the holo crystal structure were aligned to each other by the RMSD distances between their alpha carbons. The protein and ligand structures were prepared for docking, relying on the scripts included in the AutoDockTools4 distribution. <sup>72</sup> The same docking region was used for all docking calculations, which encompassed the whole protein structure and was generated by AutoDockTools4. To carry out the docking calculations, AutoDock Vina was run with the default command line options, except for the exhaustiveness option which was increased to 24, as is suggested by the authors for large docking regions and the num\_modes option which was set to twenty to obtain the twenty best poses for each ligand. The parallel execution of the docking calculations were managed with in-house scripts.

#### Ligand similarity calculations

Based on the results of the docking calculations the best ligands binding to each discovered pocket were selected. Afterwards, similarity calculations were performed between all selected ligands. These calculations employed the RDKit program package, <sup>73</sup> using its default RDKit small molecule fingerprint and the Tanimoto similarity score. <sup>74</sup>

#### Pocket description

The binding sites of the protein, discovered through computational docking calculations, were analysed with the mdpocket program, <sup>75</sup> part of the fpocket distribution. To this end, the

19 cluster representative protein structures were aligned to each other by their alpha carbons and concatenated to create a mock trajectory readable by mdpocket. The regions of space which the discovered binding pockets occupy were selected manually, by inspecting the poses of the ligands binding to the pocket in question. Based on the suggestions of the mdpocket authors, large regions were selected for each pocket, encompassing all or almost all docked ligand poses. With the protein structures concatenated and the binding regions selected, mdpocket was run with the -S option, instructing the program to score pockets by their druggability. Among its results mdpocket provides a number of pocket descriptors calculated for each frame of the supplied trajectory. From these descriptors, the pocket volume and various pocket druggability scores are utilised in the present study. To qualitatively evaluate the general druggability of a given pocket, a simple composite druggability score is defined here, that can be calculated from the descriptors provided by mdpocket as:

$$S = S_H + S_V + S_P + S_C \,. \tag{2}$$

Here,  $S_H$ ,  $S_V$ ,  $S_P$  and  $S_C$  are the hydrophobicity, volume, polarity and charge scores calculated by mdpocket respectively. It is emphasised that the definition of S is not suggested by the mdpocket developers and it is not intended as an absolute metric of pocket quality, but rather as a qualitative means to compare the druggability of the same pocket in different protein conformations. For more information about the calculation and meaning of the pocket descriptors utilised for S, see the mdpocket documentation or Reference 75.

#### Results and discussion

## Equilibration of the trajectories

In this section, the equilibration of the protein during the various (cosolvent) MD trajectories is examined by plotting the RMSD distance of the protein structure from its initial state

throughout the simulated time. On top of the usual task of selecting the equilibrated part of each trajectory to be considered for further analysis, these plots are useful to detect potential differences in the equilibration process between the traditional and the cosolvent trajectories. On Figure 1, such plots obtained for the first replica of each solvent type are shown. It is reassuring, that the protein structures seem to be well equilibrated after 50 ns of simulation time in all three cases. The equilibration appears to be happening slightly faster in the benzene and especially in the phenol cosolvent trajectory than in water. The equilibrated RMSD values plotted on Figure 1 are somewhat higher for the two cosolvent trajectories than in water. This could indicate that the cosolvent probes have stabilised some conformation is that are not often visited with water as solvent and that are farther from the original protein conformation than those appearing frequently in water based simulations.

#### Selecting representative protein conformations

As mentioned in the Computational Details, the dbscan algorithm of cpptraj is used to perform the clustering of the trajectories. The clustering is carried out separately for the three solvents, with the three replicas of each of them concatenated and treated as a single trajectory. In order to carry out a successful clustering of the trajectories, first the k and  $\varepsilon$  parameters of the density based clustering algorithm have to be tuned. On top of performing this tuning of the parameters, the effects of considering only the alpha carbon atoms for the RMSD calculations instead of all heavy atoms of the protein are also evaluated. Finally, possible redundancies in the set of representative protein structures are investigated.

The tuning of the parameters of the dbscan algorithm is performed by systematically varying the values for these parameters to see which combination yields the most optimal clustering. The computation of the RMSD distance between all frames of a trajectory is much more demanding if on top of the alpha carbons, all other heavy atoms are considered as well. To speed up these computations, the technique of sieving is utilised: only every other frame is considered explicitly during the clustering, the remaining frames are simply

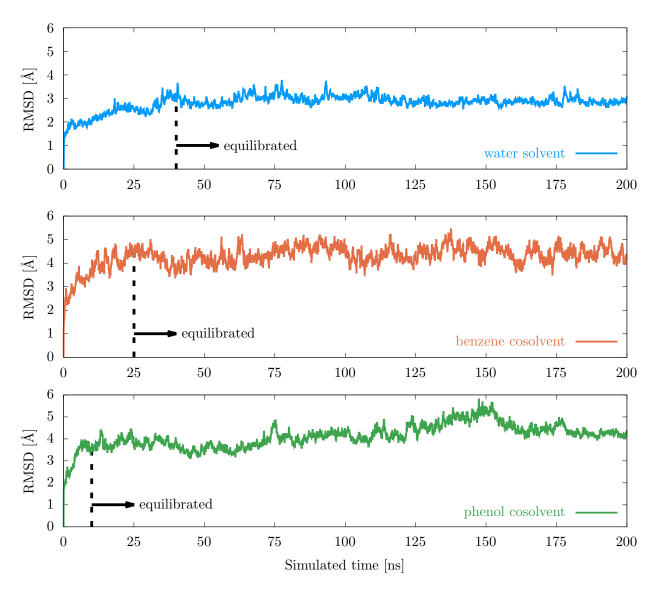


Figure 1: The evolution of the RMSD distance of the protein from the starting conformation during the MD trajectories. The first replica for each solvent is plotted.

added to the cluster with the cluster representative most similar to them. To measure the quality of the clustering four metrics are utilised. Two of these have already been discussed, namely the and the . Since both of these scores are heavily influenced by the number of obtained clusters, <sup>76</sup> the comparison of their absolute values between different MD trajectories has limited meaning. Instead, the trends arising in these metrics through the systematic variation of the clustering parameters can be interpreted to optimise these parameters. At this point it is useful to reiterate, that low values of and high values of are desirable. The other two descriptors utilised to describe the quality of the clustering are the number of noise frames (frames not included in any cluster), and the number of clusters defined by the algorithm. The number of noise frames should clearly be kept low to avoid missing any important conformations, only because it is visited very rarely and is therefore considered an outlier by the algorithm. Finally, while a high number of clusters is desirable as it can result in a wider variety of protein conformations, the computational limitations of performing explicit docking calculations to each representative conformation with thousands of ligands should be kept in mind.

On Figure 2, the descriptors of the water trajectory clustering can be seen, for the case when only the alpha carbons are considered during the RMSD distance calculations. Considering the top two plots at first, it can be observed that the and values are zero if  $\varepsilon$  is greater than or equal to 1.2 Å. The reason for this is that above this  $\varepsilon$  value all frames of the trajectory are grouped into a single cluster, for which these descriptors cannot provide a meaningful value. Since a single cluster is clearly not ideal, these large  $\varepsilon$  values do not need to be considered during the search for the optimal parameters. Focusing instead on parameter k, the most significant differences between the different values for this can be discovered on the plot. Here, the curves with k=4 or 6, reaching their peak at  $\varepsilon=1.1$ , are clearly superior to the other two. The curve corresponding to k=7 is somewhat of an outlier on this graph, with its peak at  $\varepsilon=1.15$  Å instead of 1.1 Å. On the plot, the variation of k has much more limited effects. In fact, all curves are more or less constant if  $\varepsilon$  is smaller

than or equal to 1.1 Å, at which point the values drop rapidly and become zero at 1.2 Å. Considering that  $\varepsilon=1.1$  Å is the point at which the values start decreasing, it is reasonable to assume that it is at this point that some significant changes are occurring in the way the clusters are defined. Together with the fact that  $\varepsilon=1.1$  Å provides clusterings with the best values, this observation makes this value of  $\varepsilon$  a promising candidate to be the optimal choice. By looking at the bottom two plots of Figure 2, the number of noise frames and number of clusters can be observed. On these plots one can find further advantages of the  $\varepsilon=1.1$  Å choice. These are that the number of noise frames start their rapid increase only at slightly smaller epsilon values, and that a reasonable number of clusters, thirteen, are obtained for this value.

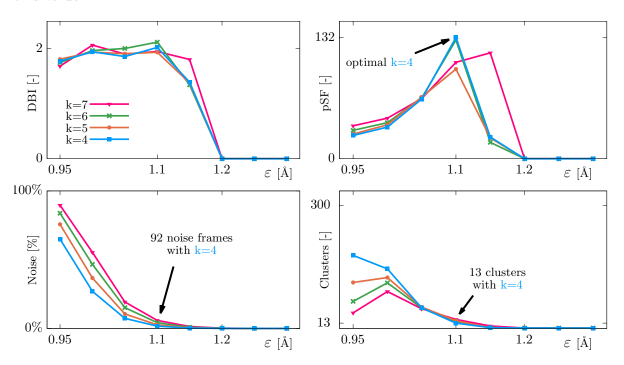


Figure 2: Plots of the clustering descriptors utilised for the tuning of the dbscan parameters. The descriptors were obtained by clustering the MD trajectory with water as the solvent and considering only the alpha carbon atoms for the RMSD calculations. The  $\varepsilon$  parameter in units of ångströms are shown on the horizontal axes in all cases, while on the vertical axes the various unitless descriptors are shown. From the top left in clockwise direction: the Davies–Bouldin index, the pseudo-F statistic, the number of clusters and the number of noise frames are shown.

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