Sensitivity in predicted relative binding free

energies from incremental ligand changes

within a model binding site

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Abstract

Despite innovations in sampling techniques for molecular dynamics (MD), reliable

prediction of protein-ligand binding free energies from MD remains a challenging prob-

lem, even in well studied model binding sites like the apolar cavity of T4 Lysozyme

L99A. In this study, we model recent experimental results that show the progressive

opening of the binding pocket in response to a series of homologous ligands. ² Even

while using enhanced sampling techniques, we demonstrate that the predicted relative

binding free energies (RBFE) are still highly sensitive to the initial protein conforma-

tional state. Particularly, we highlight the importance of sufficient sampling of protein

conformational changes and possible techniques for addressing the issue.

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1 Introduction

Medicinal chemistry programs typically focus on changes in ligand binding affinity from incremental changes to the ligand. Focus on how the protein adapts to the changes in the ligand is generally neglected. T4 L99A is well studied experimentally and computationally. It is frequently used as a model binding site in free energy prediction studies. In this study, 8 congeneric ligands were investigated, where addition of a single methyl group was used to lengthen the ligand. Through determination of protein-ligand bound x-ray crystal structures it was revealed T4 Lysozyme adopts 3 discrete conformations in response the series of growing ligands. Consideration of the protein adaptations into discrete conformations may be an important aspect in inhibitor design.

2 Methods

Protein/Ligand preparation

All proteins were prepared and aligned in Maestro³ using the 'Protein Preparation Wizard' ^{4–8} tool and with the following settings enabled (as they appear in the Maestro menu):

- Preprocess: Assign bond orders, Add hydrogens, Create zero-order bonds to metals, Create disulfide bonds, Cap termini, Delete waters beyond 5Åfrom het groups
- Refine: Sample water orientations, Use PROPKA pH: 7.0, Remove waters with less than 3 H-bonds to non-waters, and restrained minimization.

Protein structures were taken from PDBs: 4W52,4W53,4W54,4W55,4W56,4W57,4W58,4W59 corresponding to bound structures of benzene, toluene, ethylbenzene, n-propylbenzene, secbutylbenzene, n-butylbenzene, n-pentylbenzene, and n-hexylbenzene, respectively. Each simulation will start from either the protein closed state (PDB:4W52) or the open state (PDF:4W59). When using the FEP+ protocol, ligand crystal structure positions were used

as the starting position of the simulation. When using the LigandFEP protocol, a similar workflow to the tutorial⁹ was followed.

Generally, two options were taken:

- (1a) If the simulation starts from the protein closed state, the benzene crystal position was used as a reference for fragment building (PDB:4W52).
- (1b) The corresponding ligand in the transformation was built by duplicating benzene in place and adding methyl groups.
- (2a) If the simulation starts from the protein open state, the n-hexylbenzene crystal position was used as reference for fragment building (PDF:4W59).
- (2b) The corresponding ligand in the transformation was built by duplicating n-hexylbenzene in place and deleting methyl groups.

Ligand tail fragments were added using the Build/Fragments toolbar in Maestro and were not overlaid or docked. As the ligand tails were built, bonds were manually rotated so that the tail was oriented in a similar manner as in their corresponding crystal structure. Following, the newly added atoms in the tail were locally minimized, leaving the core in its initial position. This was done in an attempt to minimize the core RMSD, which LigandFEP uses to select the ligand heavy atoms to include in the REST region.

FEP protocols

Using the Schrödinger application suite (release 2015-3), ¹⁰ two FEP protocols were used in this project: FEP+¹¹ and LigandFEP. ⁹ FEP+ is a fully automated work flow that plans perturbation pathways based off the LOMAP ¹² mapping algorithm which uses the maximum common substructure (MCS) between any pair of compounds. LigandFEP is an academic toolkit that generates the configuration files to perform the free energy calculation but is limited in the sense that the user must plan each perturbation path instead. Both FEP protocols use the default Desmond relaxation protocol and the FEP/REST methodology. ^{13–16}

Results and discussion sections will present information from using the LigandFEP protocol, additional data not discussed here can be found in the supplementary info.

REST region selection

In this study, by default, only heavy atoms in the ligand were included in the REST region unless specified otherwise. Further details on the temperature profile and how the REST region is normally selected can be found in previous studies ^{11,15} in the supporting information. Simulations that included protein heavy atoms in the REST region are referred to with the 'pREST' label, where selection of the particular residues is described as follows. Based on visual inspection of our molecular dynamics simulations and considering the F-helix spans residues 107-115, we selected residues Glu108, Val111, and Gly113 to include into the REST region (Fig 10). Glu108 sits near the start of the helix which appears as a hinge point for the opening and closing of the binding cavity (Fig 11). Following, Val111 appears in the middle of the helix and was observed to undergo the largest motion during protein conformational changes (Fig 12). Gly113 was included in order to collectively have hot regions approximately at the start, middle and end points of the helix.

Simulation details

Desmond ^{17–20} simulation protocols have been described previously in the supporting information ¹¹ or can be found in greater detail in the Desmond User Manual. ²¹ In summary, solute molecules are restrained to their initial positions while minimizing using a Brownian dynamics NVT integrator for 100ps, followed by 12ps simulations at 10K with a NVT ensemble and then a NPT ensemble using the Langevin method. ²² Next, a 24ps simulation followed by a final 240ps simulation with solute molecules unrestrained, both are carried at room temperature with a NPT ensemble using Langevin. Production simulations were carried out to a length of up to 55ns for closed-open transformations and up to 25ns for closed-intermediate transformations. FEP/REST simulations were run on four GeForce

GTX Titan Black GPUs using the Desmond/GPU engine with OPLS2005²³ and OPLS3²⁴ forcefield parameters. Calculated free energies were determined using the Bennett acceptance ratio²⁵ (BAR) with error estimations using both bootstrapping and BAR analytical error prediction.²⁶ Hysteresis around closed thermodynamic cycles and best estimates of the free energies with their errors were calculated using the cycle closure algorithm discussed in a previous publication.¹⁵

RMSD analysis

3 Results

Calculated free energies depend strongly on starting protein conformation

Using the default FEP/REST methodology, ¹⁵ we find calculated free energies significantly depend on the protein starting conformation, especially for large perturbations (i.e. opening the cavity from the closed state). To illustrate this, we begin our molecular dynamics simulations both from the protein closed and open conformations then perform alchemical transformations to ligands that occupy the opposite protein conformational state. In this study, root-mean-square-deviation (RMSD) of the backbone atoms in the F-helix is used to determine the conformational state of the protein over the course of the simulation. Here, we demonstrate the default 5ns simulation time and REST region selection in the Schrödinger FEP workflow are insufficient for adequate sampling of the motion in the F-helix and does not eliminate the dependence on the initial protein state.

An examination of the largest alchemical transformation, benzene to n-hexylbenzene, highlights the sampling problems faced when using the standard FEP/REST protocol. From experimental data of ligand occupancies (Table 1), we expect in our simulations of n-hexylbenzene to see the protein primarily in the open state over the closed state. Instead,

we find the protein remains trapped in its initial conformational state whether we start from closed (Fig 1) or open (Fig 2) over the course of the 5ns simulation. From the protein closed simulation, the protein only begins to enter the intermediate state around 3ns but never enters the open conformation. As the protein tries to accommodate n-hexylbenzene and enter its preferred open state, protein-ligand strain results, yielding a positive value for $\Delta\Delta G_{calc}(+4.13 \text{ kcal/mol})$. On the other hand, in the protein open simulations, the protein already begins in its preferred state for n-hexylbenzene and stays only in this open state. As expected, the $\Delta\Delta G_{calc}$ comes out negative(-0.61 kcal/mol) as there is no occurrence of large protein-ligand strain in order to open the cavity. Ultimately, we arrive at two very different relative free energies values, where the discrepancy is as large as +4.74 kcal/mol for the same transformation of benzene to n-hexylbenzene¹. Collectively, when we view the discrepancy of all calculations involving closed-open transformations we find the root-mean-square-inconsistency (RMSI) to be +4 kcal/mol (Table 6). Clearly, despite the use of FEP/REST, we are unable to adequately sample all the relevant states within the standard 5ns time frame, resulting in such large differences in $\Delta\Delta G_{calc}$.

In the case of more moderate alchemical transformations, such as cases that involve the set of closed ligands (i.e. benzene to n-propylbenzene) to intermediate ligands (i.e. sec-/n-butylbenzene), we find that the calculated free energies still have some (albeit much smaller) dependence on the initial protein conformation using the default protocol. For the set of transformations to the intermediate state, the RMSI in $\Delta\Delta G_{calc}$ for protein closed versus open simulations is +0.60 kcal/mol (Table 3). However, when we compare $\Delta\Delta G_{calc}$ against $\Delta\Delta G_{exp}$ for transformations involving n-butylbenzene, we find that simulations starting from the protein closed conformation are further from converging to $\Delta\Delta G_{exp}$ than when starting from the protein open conformation. From the protein closed simulations, the RMS error relative to $\Delta\Delta G_{exp}$ is +1.40 kcal/mol (Table 9), while for the protein open

¹It should be noted that the binding affinities of n-pentyl/n-hexylbenzene to T4-L99A are not known and were inaccessible in experimental studies due to solubility limits. ² For cases involving these ligands, we only focus on the convergence of the calculated free energies between simulations starting from protein closed or open.

simulations it is +0.70 kcal/mol (Table 10). Based on experimental evidence (Table 1), we should expect to see some sampling (30%) of the open conformation for the n-butylbenzene ligand if the calculations are converged. Evidently, we find the simulations remain trapped in their respective starting conformations, resulting in inadequate sampling in the protein closed simulations (Fig. 3) versus the protein open simulations (Fig. 4). Despite performing much smaller alchemical transformations, we still encounter sampling problems that result in $\Delta\Delta G_{calc}$ that depend on the intial protein conformation, evident when comparing to $\Delta\Delta G_{exp}$.

Including protein residues into the REST region (pREST) improves sampling

Primarily, we encounter major sampling problems when we begin our simulations from the protein closed state and attempt a mutation which should result in the helix opening. In order to facilitate protein motion, we included 3 key residues spanning the F-helix region into the REST region, which we will denote simulations using this with pREST. By expanding the REST region, we are able to drive the F-helix out its initial state trap faster by locally heating up key regions and thereby reduce our problem with inadequate sampling.

To demonstrate the REST improvement over the default protocol, we return to the case of benzene to n-hexylbenzene. Here, we show the facilitation of the helix motion by first referring to Fig 1 which shows that there is no sampling of the open state for the default protocol. On the other hand with the pREST, we see a few open state points around 3ns and even a single open point before closing again in our initial step. (Fig 5. Alternatively, we can further illustrate the enhancement of protein transitions by viewing all replicas collectively. In Fig 6 and Fig 7, we perform the same RMSD analysis but instead represent each time point as a colored bar and no longer plot the raw RMSD. Visually, it is easy to see that there are far less transitions in default simulations (Fig 6) as opposed to pREST simulations (Fig 7).

Collectively, for all our closed/open and closed/intermediate transformations using pREST, we find only some minor improvements in the RMSI and even cases where we perform worse. For closed/open transformations (Table 7), the RMSI improves to +2.78 kcal/mol (previously +4 kcal/mol) but gets regresses slightly to +0.78 kcal/mol (previously +0.60 kcal/mol) for closed/intermediate cases (Table 4). In general, simulations starting from the closed state had $\Delta\Delta G_{calc}$ values that moved towards favorability (i.e. more negative $\Delta\Delta G_{calc}$) and while protein open simulations $\Delta\Delta G_{calc}$ values tended towards unfavorability (i.e. more positive $\Delta\Delta G_{calc}$). This is indicative of the fact that pREST is indeed improving sampling, but it is evident that our $\Delta\Delta G_{calc}$ are far from convergence given the RMSI is still large, especially for closed/open transformations.

Long simulations enhances protein conformational sampling from more exchanges

Although we see improvements in sampling with pREST, the standard implemented time frame of 5ns clearly is not long enough completely capture the transition of closed to open in the helix. Here, we simulate 55ns for closed/open transformations and take the final 15ns of the simulation while for closed/intermediate we simulate 25ns and take the last 10ns of the simulation, discarding the initial time as additional equilibration time. In simply running longer, we allow our simulations to perform more exchanges across replicas and thereby allow for better sampling of all conformational states.

Returning to our most extreme transformation, benzene to n-hexylbenzene, we have shown pREST alone does not allow for adequate sampling of the open state (Fig 5). Now, when we run much longer we see far more sampling of the open protein conformational state in the final 10ns window (Fig 8). In viewing all the replicas (Fig 9), we illustrate the dramatic increase in protein conformational sampling in stark contrast to our 5ns simulations (Fig 7).

By simulating longer with pREST we dramatically increase our sampling of the intermediate/open states and almost entirely eliminate the dependence on the initial protein conformational state. For the set of closed/open transformations the RMSI dramatically falls to +0.57 kcal/mol (Table 8). In the set of closed/intermediate transformations, only a few cases required an extended simulation time of 25ns, after doing so we obtain an overall RMSI of +0.42 kcal/mol (Table 5). There still remains some discrepancy between $\Delta\Delta G_{calc}$ from protein open or closed simulations, but it now falls within a much more reasonable range of less than +1 kcal/mol.

4 Discussion

In this study, we find that relative free energy calculations can suffer from substantial convergence problems resulting from relatively modest protein conformational changes. Although, the protein conformational changes in T4 lysozyme (L99A) are extremely localized to a rearrangement of a single helix, we still encounter sampling challenges. These problems have profound implications for the accuracy of computed relative free energies in these cases. Particularly, we find that calculated relative free energies can depend on the initial protein conformational state by up to 4 kcal/mol.

By looking at cases that involve a conformation change in the protein, we show the $\Delta\Delta G_{calc}$ is sensitive to the initial protein conformational state. Such cases are when the alchemical transformation involves mutating ligands that primarily occupy the closed state (i.e. benzene to n-propyl) into ligands that occupy the intermediate (i.e. sec-/n-butyl) or open states (n-pentyl/n-hexyl) (Table 1). In tracking the RMSD relative to the crystallographic protein closed structure, we show the protein remains trapped in its initial state throughout the simulation when using the implemented default protocol. Through remaining trapped, we are unable to adequately sample the correct protein-ligand conformational states and thereby poorly computed $\Delta\Delta G_{calc}$.

Without prior knowledge of preferred protein conformational states on ligand binding, we can arrive at very different binding affinity predictions that are sensitive to the initial protein state. Generally, from protein closed simulations we found $\Delta\Delta G_{calc}$ were highly positive and unfavorable, a result from strain in the protein as the ligand begins to grow in the binding cavity. However, in protein open simulations we tended to see $\Delta\Delta G_{calc}$ that were negative and favorable as no protein-ligand strain is encountered (Table 6). If we only had the crystal structure of the closed protein-ligand complexes, we would blindly conclude that the much larger, open ligands bind worse than smaller ones. On the other hand, if only the open protein-ligand complexes were available, the opposite would be concluded in that larger ligands are better binders than smaller ligands.

By including key residues into the REST region and simulating longer, we reduce the $\Delta\Delta G_{calc}$ dependence on the initial protein conformation to a more reasonable range of less than 1 kcal/mol. Through expanding the REST region, intermediate lambda windows are able to more easily access the intermediate and open conformations by effectively heating key residues that would facilitate protein motion, illustrated in Fig 7. Further, by simulating longer we allow for more exchanges between replicas, which in turn enhances sampling at our physically relevant end state lambda windows (Fig 9). With these modifications to the default protocol, we almost completely converge our $\Delta\Delta G_{calc}$ to the same value regardless of the starting protein conformation.

From our results, we present strong evidence for the importance of studying kinetically distinct protein conformational states prior to performing binding free energy calculations. Without performing such initial studies, early drug discovery projects are faced with the hazard that the effect of the initial protein structure on predicted binding free energies is essentially hidden. Only from prior crystallography studies² and our systematic trials of varying the protein starting structure were we able to identify this problem and correct it. Generally, our brute-force approach of simulating longer and multiple trials with varied protein structures is not a desirable or a even a feasible approach, especially in early drug discovery phases. At the industrial level, ligand libraries can be large—driving computational cost exponentially if we simulate longer—or experimental structures can be sparse

for new therapeutic protein targets. For future studies, approaches using Markov State Models (MSMs)²⁷ can potentially be of great use for identifying discrete protein conformations. MSMs build a representation of the conformational space from batches of short molecular simulations, whereby the discrete states and transition rates between them can be determined. Utilizing MSMs can thereby provide useful insight on the various protein conformational states before running free energy predictions.

5 Conclusions

Overall, we have shown that the presence of kinetically distinct protein conformational states—that are modulated by ligand binding—can dramatically impact the accuracy free energy calculations. Had we not known of the various protein-ligand states, we would not have been able to see our predictions had a strong dependence on the protein-ligand structure we had started our simulations from. In this study, we would have encountered a worst-case scenario if we only had the structure of the apo protein available, as is often the case in early stages of drug discovery. Then, a medicinal chemist could be tasked with docking a library of ligands and running binding free energy predictions in order to determine the most suitable candidates to push forward. Dangerously, the chemist would then discard ligands with apparently low binding affinities, where these ligands only appear unfavorable because of unsampled protein conformational changes. Without prior knowledge of the existence of other protein conformational states, one would never know that these calculations are actually incorrect due to poor convergence from insufficient sampling.

Although FEP calculations have shown tremendous recent successes, ¹¹ we are still faced with challenges in adequate protein sampling. Using T4 lysozyme (L99A) as our model system, we highlight sampling problems even from a relatively small (1-3Å) and localized single helix rearrangement in response to a series of growing ligands. In this study, we show that using a typical 5ns simulation time with only ligand atoms in the REST region leads to

free energies that significantly depend on the initial protein conformational state. Only by longer simulation times and expansion of the REST region to include key protein residues were we able to eliminate the dependence on the starting conformation and come close to convergence. More importantly, this demonstrates that special attention and care should be exercised when performing binding affinity predictions where regions of flexibility surround the binding site.

Figures

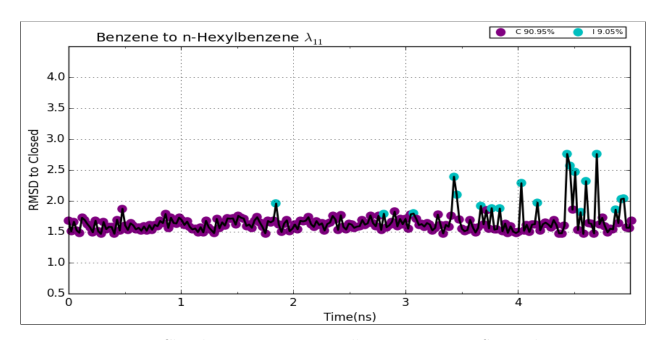


Figure 1: Closed - Benzene to n-Hexylbenzene 0-5ns RMSD Replica11

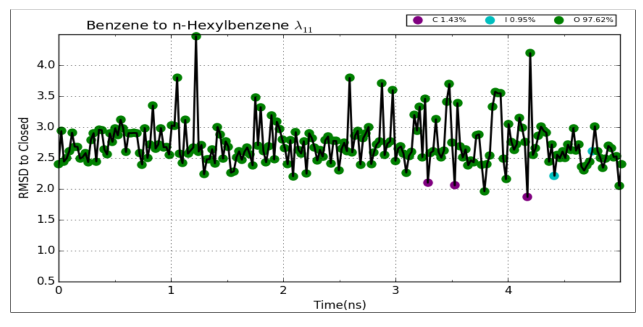


Figure 2: Open - Benzene to n-Hexylbenzene 0-5
ns RMSD Replica 11

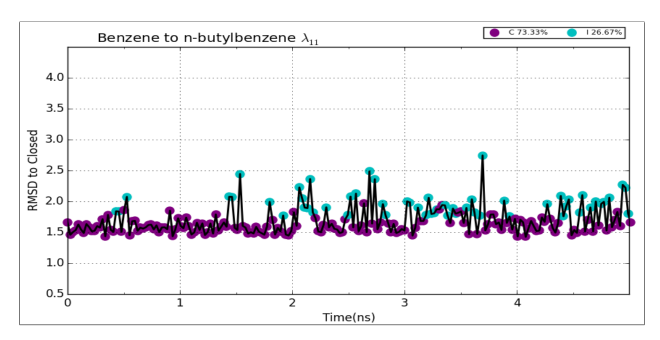


Figure 3: Closed - Benzene to n-butylbenzene 0-5ns RMSD Replica11

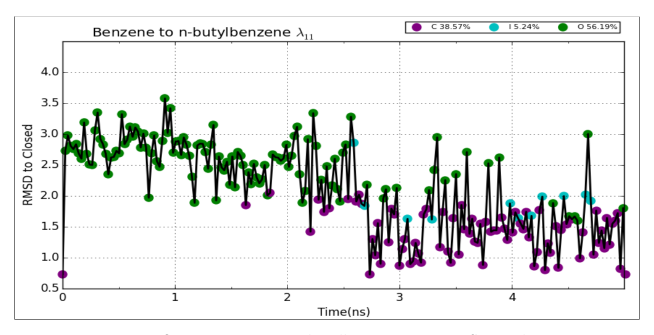


Figure 4: Open - Benzene to n-butylbenzene 0-5
ns RMSD Replica 11

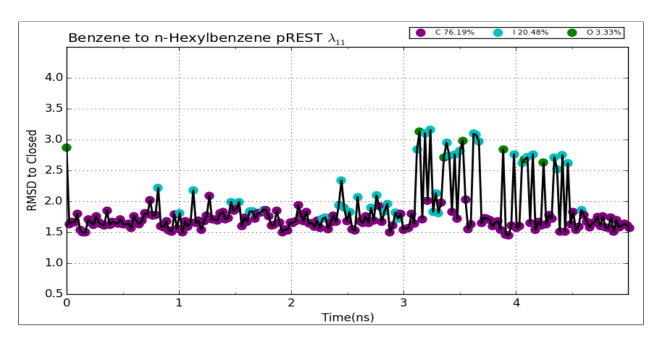


Figure 5: Closed - Benzene to n-Hexylbenzene 0-5ns RMSD Replica11 pREST

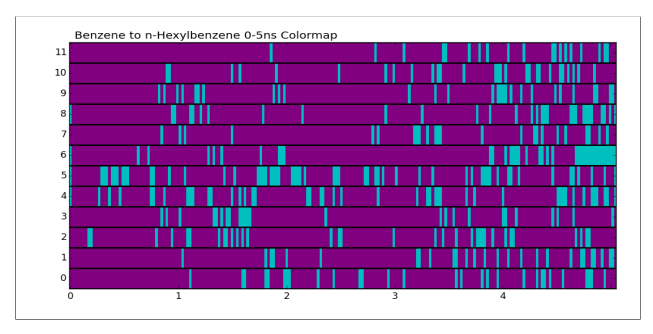


Figure 6: Closed - Benzene to n-Hexylbenzene 0-5ns Colormap

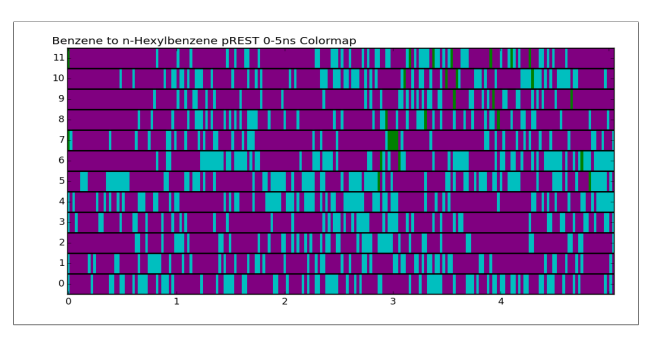


Figure 7: Closed - Benzene to n-Hexylbenzene 0-5ns Colormap pREST

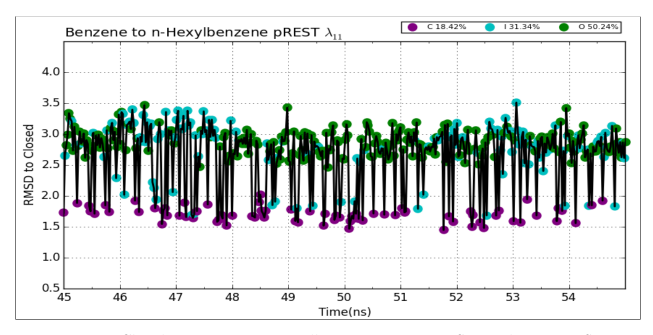


Figure 8: Closed - Benzene to n-Hexylbenzene 45-55ns RMSD Replica11 pREST

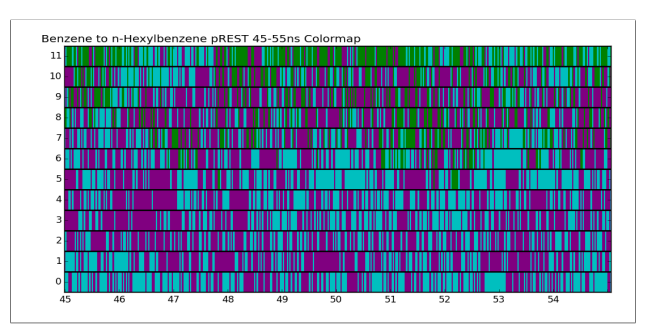


Figure 9: Closed - Benzene to n-Hexylbenzene 45-55ns Colormap pREST

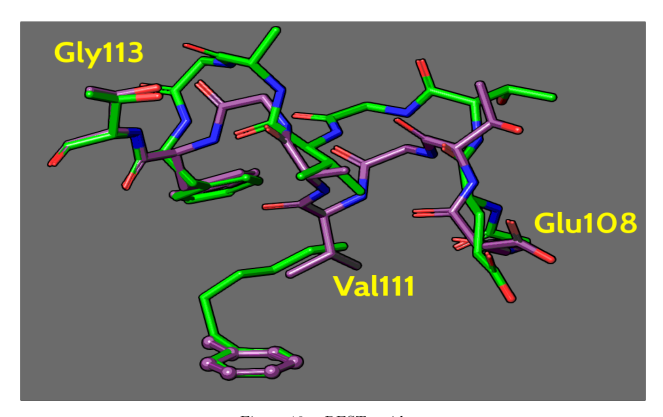


Figure 10: pREST residues

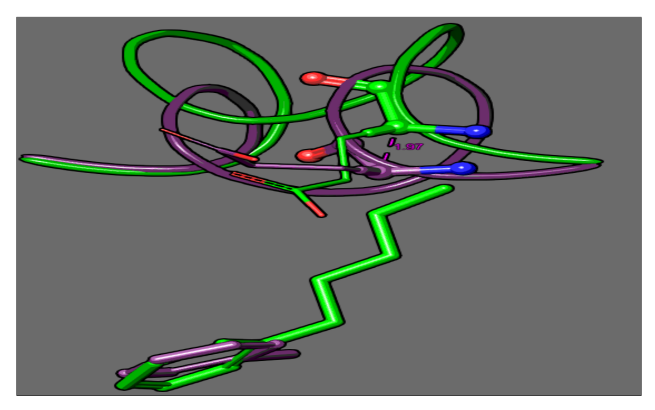


Figure 11: Residue Glu108

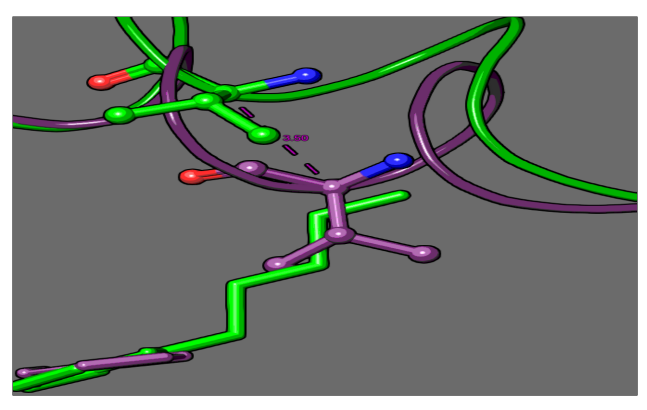


Figure 12: Residue Val111

Tables

Table 1: Loop Occupancies 2

Ligand	С	I	О
benzene	0.9	-	-
toluene	0.8	0.2	-
ethylbenzene	0.5	0.5	-
n-propylbenzene	0.6	0.4	-
sec-butylbenzene	0.4	0.6	-
n-butylbenzene	0.1	0.6	0.3
n-pentylbenzene	0.3	-	0.7
n-hexylbenzene	0.3	-	0.7

Table 2: Ligand Binding Affinities

PDB	Ligand	ΔG_{exp}	Error
4W52	benzene	-5.19	0.16
4W53	toluene	-5.52	0.04
4W54	ethylbenzene	-5.76	0.07
4W55	n-propylbenzene	-6.55	0.02
4W56	sec-butylbenzene	N/A	-
4W57	n-butylbenezene	-6.70	0.02
4W58	n-pentylbenzene	N/A	-
4W59	n-hexylbenzene	N/A	-

Table 3: Closed-Intermediate Transformations

Ligand 1	Ligand 2	Closed	Err	Open	Err	C-O Diff
benzene	n-butylbenzene	0.58	0.07	-0.59	0.09	1.17
toluene	n-butylbenzene	-0.28	0.06	-1.27	0.09	0.99
ethylbenzene	n-butylbenzene	0.24	0.07	-0.23	0.07	0.47
n-propylbenzene	n-butylbenzene	0.99	0.06	0.63	0.04	0.36
benzene	sec-butylbenzene	2.36	0.09	2.14	0.11	0.22
toluene	sec-butylbenzene	1.47	0.07	1.14	0.09	0.33
ethylbenzene	sec-butylbenzene	1.90	0.08	1.77	0.07	0.13
n-propylbenzene	sec-butylbenzene	2.86	0.06	2.67	0.05	0.19

Table 4: Closed-Intermediate Transformations pREST

Ligand 1	Ligand 2	Closed	Err	Open	Err	C-O Diff
benzene	n-butylbenzene	-0.10	0.11	-0.72	0.12	0.62
toluene	n-butylbenzene	0.90	0.09	-0.36	0.09	1.26
ethylbenzene	n-butylbenzene	-0.20	0.09	-0.43	0.08	0.23
n-propylbenzene	n-butylbenzene	1.00	0.07	0.49	0.06	0.51
benzene	sec-butylbenzene	0.45	0.08	1.34	0.12	0.89
toluene	sec-butylbenzene	0.60	0.12	0.60	0.10	0.0
ethylbenzene	sec-butylbenzene	1.09	0.10	1.69	0.09	0.60
n-propylbenzene	sec-butylbenzene	1.88	0.07	3.05	0.08	1.17

Table 5: Closed-Intermediate Transformations pREST 15-25ns

Ligand 1	Ligand 2	Closed	Err	Open	Err	C-O Diff
benzene	n-butylbenzene	-1.15	0.09	-0.72	0.12	0.43
toluene	n-butylbenzene	-0.52	0.09	-0.36	0.09	0.16
ethylbenzene	n-butylbenzene	-0.20	0.09	-0.43	0.08	0.23
n-propylbenzene	n-butylbenzene	0.53	0.07	0.49	0.06	0.04
benzene	sec-butylbenzene	0.45	0.08	1.34	0.12	0.89
toluene	sec-butylbenzene	0.60	0.12	0.60	0.10	0.0
ethylbenzene	sec-butylbenzene	1.09	0.10	1.69	0.09	0.60
n-propylbenzene	sec-butylbenzene	1.88	0.07	1.84	0.06	0.04

Table 6: Closed-Open Transformations

Ligand 1	Ligand 2	Closed	Err	Open	Err	C-O Diff
benzene	n-pentylbenzene	2.36	0.12	-1.33	0.11	3.69
toluene	n-pentylbenzene	1.77	0.09	0.34	0.10	1.43
ethylbenzene	n-pentylbenzene	2.45	0.08	0.46	0.09	1.99
n-propylbenzene	n-pentylbenzene	3.46	0.08	-0.22	0.08	3.68
benzene	n-hexylbenzene	4.13	0.16	-0.61	0.15	4.74
toluene	n-hexylbenzene	2.90	0.14	-1.63	0.08	4.53
ethylbenzene	n-hexylbenzene	3.63	0.11	-0.76	0.09	4.39
n-propylbenzene	n-hexylbenzene	5.85	0.10	0.13	0.06	5.72

^a Some text; ^b Some more text.

Table 7: Closed-Open Transformations pREST

Ligand 1	Ligand 2	Closed	Err	Open	Err	C-O Diff
benzene	n-pentylbenzene	1.45	0.13	0.15	0.10	1.30
toluene	n-pentylbenzene	1.40	0.13	0.82	0.11	0.58
ethylbenzene	n-pentylbenzene	2.89	0.10	1.32	0.10	1.57
n-propylbenzene	n-pentylbenzene	4.40	0.12	1.06	0.09	3.34
benzene	n-hexylbenzene	2.74	0.19	1.37	0.13	1.37
toluene	n-hexylbenzene	3.21	0.15	-1.08	0.09	4.29
ethylbenzene	n-hexylbenzene	3.39	0.11	-0.14	0.10	3.53
n-propylbenzene	n-hexylbenzene	4.93	0.12	1.28	0.10	3.65

^a Some text; ^b Some more text.

Table 8: Closed-Open Transformations pREST 40-55ns

Ligand 1	Ligand 2	Closed	Err	Open	Err	C-O Diff
benzene	n-pentylbenzene	1.86	0.06	1.50	0.06	0.36
toluene	n-pentylbenzene	1.03	0.06	0.71	0.06	0.32
ethylbenzene	n-pentylbenzene	1.69	0.06	1.60	0.06	0.09
n-propylbenzene	n-pentylbenzene	3.43	0.04	2.44	0.04	0.99
benzene	n-hexylbenzene	2.14	0.08	1.41	0.07	0.73
toluene	n-hexylbenzene	0.33	0.08	1.16	0.06	0.84
ethylbenzene	n-hexylbenzene	1.97	0.07	2.39	0.06	0.42
n-propylbenzene	n-hexylbenzene	3.49	0.06	3.44	0.05	0.05

^a Some text; ^b Some more text.

Table 9: Closed to n-butylbenzene Transformations (Closed Protein)

Ligand 1	Ligand 2	Closed	Err	C-Exp Err
benzene	n-butylbenzene	0.58	0.07	2.09
toluene	n-butylbenzene	-0.28	0.06	0.90
ethylbenzene	n-butylbenzene	0.24	0.07	1.18
n-propylbenzene	n-butylbenzene	0.99	0.06	1.14

Table 10: Closed to n-butylbenzene Transformations (Open Protein)

Ligand 1	Ligand 2	Open	Err	O-Exp Err
benzene	n-butylbenzene	-0.59	0.09	0.92
toluene	n-butylbenzene	-1.27	0.09	0.09
ethylbenzene	n-butylbenzene	-0.23	0.07	0.71
n-propylbenzene	n-butylbenzene	0.63	0.04	0.78

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Supporting Information Available

6 Experimental

6.1 Discrete Conformations and the Ligands

T4 L99A contains an engineered apolar cavity which is our binding site of interest. The 8 congeneric ligands are all apolar and begins with a simple benzene ring. Subsequent ligands are simply addition of a methyl group to generate a growing tail up until n-hexylbenzene. In response to the growing ligand, the crystal structures show the protein will adopt into 3 conformations aptly named: closed, intermediate, and open. Primarily, the motion of the protein occurs in the F-helix (residues 107-115), which serves as a sort of gating mechanism into the apolar cavity. As the ligand tail expands, the F-helix transitions from closed to open, exposing the cavity to the bulk solvent. From observed electron densities of the F-helix in²-Fig2, the ligands occupy each of the conformations given in Table: 1

From a protein conformation clustering analysis, shown, ² Val111 occupies 3 distinct states iin accordance to the 3 protein conformations. Visualization of side-chain Val111 from the x-stal structures, shows the backbone alpha-carbon moving 1.25Åwhen transitioning from closed to intermediate, 3.25Åintermediate to open, and 3.50Åclosed to open. From our MD simulations, it is primarily the repulsive interactions between the ligands and Val111 that drive the F-helix to the open state.

6.2 Ligand Binding Affinities

Table 2 Details of the ligand binding affinities and how they were obtained can be found in the paper²

Section "Energy of Ligand Binding and Conformational Strain"

Ligands n-pentylbenzene and n-hexylbenzene affinities are inaccessible due to solubility limits. Generally, as the ligands grow from benzene to n-butylbenzene, the affinity rises linearly.

7 FEP+ vs. LigandFEP

- Comparisons between the two protocols will be used to show that, although limited,
 LigandFEP does not result in a difference in performance of accurate predicted relative
 binding free energies.
- Comparisons between FFs will show that the new and improved OPLS3 parameters are give better RBFE predictions for larger transformations.
- ***INSERT TABLE COMPARING OPLS2005/OPLS3 BETWEEN TWO PROTO-COLS***

Both are within the same MUE/RMSE range

- Highlight inconsistency in predicted RBFE when the protein starting conformation is varied.
- Predicted ddGs are in the wrong direction frequently when starting from protein closed.

 If we assume smaller to larger ligands should yield favorable (-) ddGs.

8 OPLS2005 vs. OPLS3

9 Case studies

9.1 Small Ligands

- Small transformations and ligands generally occupy protein closed state.
- Equal performance with either FF.
- Highlight good case: Toluene to Ethylbenzene with OPLS3
 - Small perturbation: Adding one carbon

- Both ligands occupy the closed state with some intermediate
- Protein starting conformation does not result in a large discrepancy in predicted ddGs.
- ***INSERT RMSD GRAPH***
- RMSD graph over the lambda=0 and lambda=1 corresponding to the toluene and ethylbenzene end states shows at each time point which conformation (reference to crystal structure) the simulation has the lowest RMSD to.
 - Purple = Closed, Teal = Intermediate, and Green = Open.
- Highlight good sampling/number of transitions between either state when starting from closed.
- Highlight points 0-1ns are still stuck in open conformation. 240ps relaxation protocol wasn't sufficient to discard few open points.
 - Show that this does not have a large impact in the final ddG from sliding time.
- Highlight not-so-good case: Toluene to n-propylbenzene
 - Perturbation involves adding two carbons
 - Discrepancy from protein starting conformation increases.
 - Highlight more open points are present (0-2ns) in open runs.
- Growing ligands require more time for helix to relax out of open state.
- Inclusion of protein residues in REST region should allow for faster transition between states which will give us a lower discrepancy between open/closed runs.
- Apply pREST to previous case and show faster relaxation out of open conformation resulting in a lower discrepancy.
- ***INSERT COMPARISON BETWEEN NORMAL AND PREST RUNS WITH EX-PERIMENTAL ddG ***

• Show pREST increases agreement with experiment and lowers error from protein starting conformation.

9.2 Intermediate Ligands

Insert all data

9.3 Open Ligands

Simulations starting from closed give a predicted ddG of +2.74 kcal/mol and from open +1.37 kcal/mol. Upon closer inspection of the closed simulations corresponding to the final state of n-hexylbenzene, we find that the protein does not remain trapped in its initial state. Instead, the region of the helix around Gly113 briefly opens to relieve the strain but quickly closes back. Hence, we still see some strain energy as the protein fails to completely stabilize into the open conformation. Viewing the open simulations, reveals that the protein is no longer trapped only in the open state and makes transitions into the intermediate and closed states. In making these transitions, the protein also experiences strain as the tail pushes the F-helix back into open from the other conformational states.

INSERT RMSD GRAPH SHOWING TRANSITIONS BETWEEN STATES FOR BENZENE TO NHEXYL

By comparing these graphs we show that inclusion of protein residues in the REST region does allow for more/faster transitions between states

***Compare pREST runs with default REST in order to show that pREST allows for the protein to get of being trapped in its initial state.

Here we have shown pREST allows for more motion in the helix in comparison to the default REST protocol. Although there is a reduction in the discrepancy, it is still fairly large (+1.37 kcal/mol). Thus, we cannot say these are converged, where this poor conver-

gence results from inadequate sampling of all the protein conformational states. Such is the case of closed pREST simulations of n-hexylbenzene, where we only see a partial opening of the helix. Similarly, in open pREST simulations with n-hexylbenzene, the helix does not enter the closed conformation. Where as it is expected to at least partially occupy the closed state, according to the loop/ligand occupancy table. It seems likely that the overall motion of the helix is not entirely accessible in the range of up to 50ns (longest we have simulated).

INSERT DATA FROM 55ns pREST simulations

Table showing all closed to open cases

Here we can show that the discrepancy does not entirely go away, but gradually gets smaller with smaller perturbations

Show that in 50ns the protein is only barely able to stabilize in the open state and vice versa

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