

Enhanced sampling approach to the induced-fit docking problem in protein-ligand binding: the Case of mono-ADP-ribosylation hydrolase inhibitors

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Enhanced Sampling Approach to the Induced-Fit Docking Problem in Protein–Ligand Binding: The Case of Mono-ADP-Ribosylation Hydrolase Inhibitors

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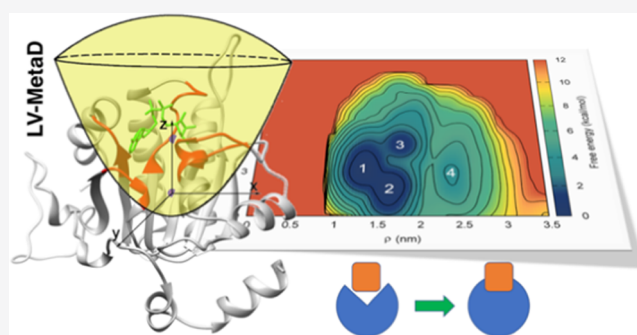


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ABSTRACT: Enhanced sampling methods can predict free-energy landscapes associated with protein/ligand binding, characterizing the involved intermolecular interactions in a precise way. However, these *in silico* approaches can be challenged by induced-fit effects. Here, we present a variant of volume-based metadynamics tailored to tackle this problem in a general and efficient way. The validity of the approach is established by applying it to substrate/enzyme complexes of pharmacological relevance: mono-ADP-ribose (ADPr) in complex with mono-ADP-ribosylation hydrolases (MacroD1 and MacroD2), where induced-fit phenomena are known to be significant. The calculated binding free energies are consistent with experiments, with an absolute error smaller than 0.5 kcal/mol. Our simulations reveal that in all circumstances, the active loops, delimiting the boundaries of the binding site, undergo significant conformation rearrangements upon ligand binding. The calculations further provide, for the first time, the molecular basis of ADPr specificity and the relative changes in its experimental binding affinity on passing from MacroD1 to MacroD2 and all its mutants. Our study paves the way to the quantitative description of induced-fit events in molecular recognition.



INTRODUCTION

Structure-based drug design (SBDD) analyzes at the atomic level how ligands interact with their biological targets. It identifies new ligands by mapping biological activity data to distinct structural and energetic features of ligand–target complexes. *In silico* predictions of the poses and affinities of ligands for their biological targets constitute a key step of the approach.^{1,2}

The predictions challenge the computational chemist when considerable structural adaptations of the ligand and the target are involved.^{3,4} Indeed, induced-fit mechanisms play a role beyond the better ligand juxtapositioning in the binding site: slow targets' conformational transitions induced by ligands during the binding/unbinding process govern the differences of the free energies of all states along the binding/unbinding pathway.^{5,6} These differences can determine ligand specificity even in structurally highly similar targets, with changes in affinity up to several 100-fold.⁵ Thus, the knowledge of these pathways can reveal induced-fit mechanisms and allow the calculation of other important pharmaceutical parameters like residence time.^{7–11}

Traditionally, in a drug design setting, medicinal and computational chemists use molecular docking that has the enormous advantage of being computationally cheap.¹² This approach allows screening poses and trends in affinities for

10⁶–10⁷ compound libraries in the order of days (with performances depending on docking software and rotational freedom of the involved compounds¹³). However, these approaches quantify target–ligand energies in terms of simple scoring functions,^{14–16} which generally neglect entropic contributions of the binding along with the solvent effects.¹⁷ As a result, scoring functions can be used at most for qualitative comparisons. In addition, and most importantly for the present discussion, they cannot predict (un)binding pathways and oversimplify the binding process: usually, only residues around the ligand are flexible, under the assumption—not justifiable with induced fit—that neither the binding site nor the binding mode is changed significantly. Therefore, docking algorithm performance drops in the presence of induced fit even in predicting poses and affinity trends. These limitations can in theory be overcome by predicting (un)-binding processes and affinities by means of molecular dynamics (MD) simulations. The main issue in this approach

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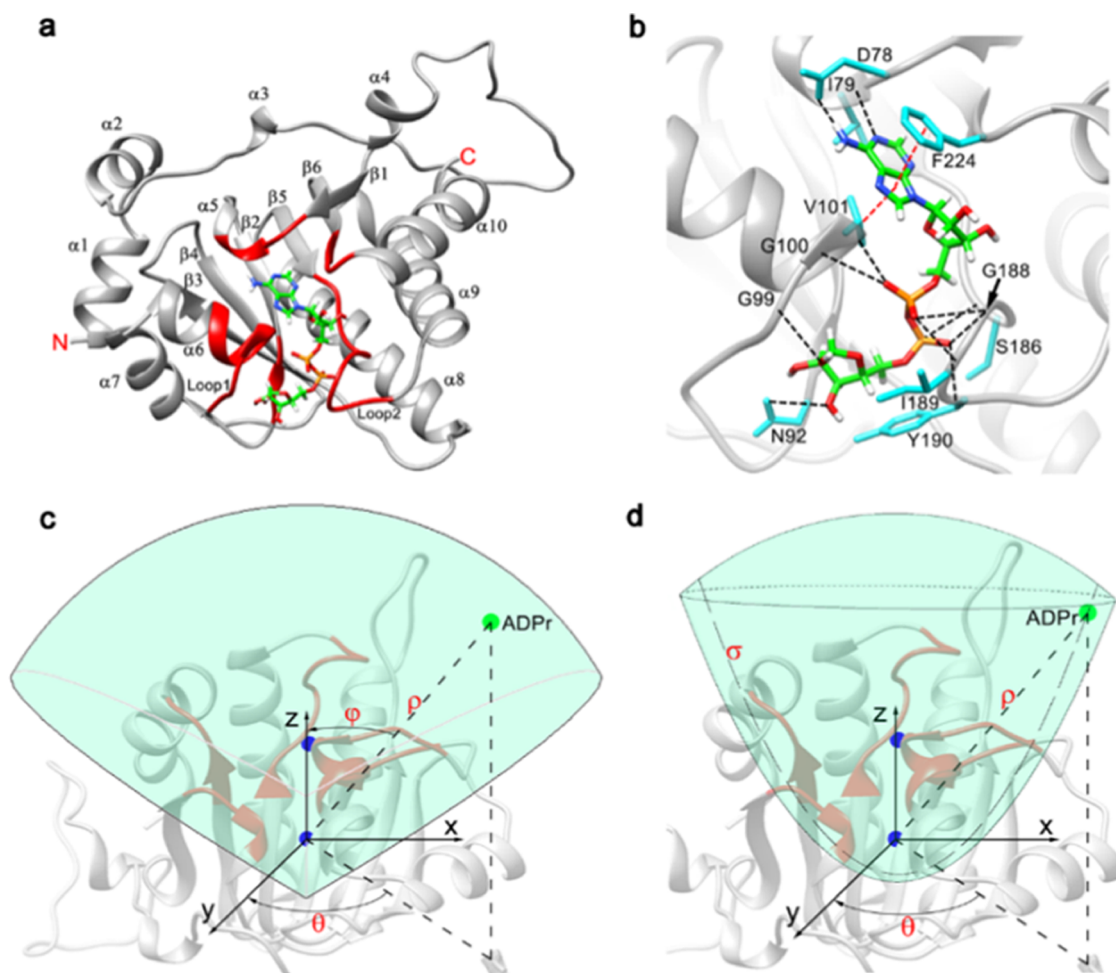


Figure 1. MacroD2 X-ray structure and the volume used in our localized volume-based metadynamics simulations. (a) WT holo MacroD2 X-ray structure (PDB code: 4IQY). The protein features a three-layered α - β - α sandwich-like conformation. The ligand ADPr is represented as sticks. The part of the protein involved in the binding pocket is highlighted in red. Loop1, 97-GGGGV-101, and loop2, 188-GIYG-191, are highlighted in red color. The backbone of the parts in gray undergoes a light constraint. (b) ADPr pose. Hydrogen bonds and hydrophobic interactions are depicted as black and red dashed lines, respectively. (c) One-third sphere-solid restraint volume. The ρ , θ , and φ spherical coordinates define the ADPr position. (d) Parabolic-solid restraint potential. The ρ , θ , and σ coordinates identify the position of the restraint potential. In both (c) and (d), the residues involved in the ligand-binding site (in red) are completely contained in the restraining volume. The center of mass of the protein and binding site (shown in blue balls) are considered to define the orientation of the volume.

is given by the timescales in which these processes live (from the order of the microsecond and longer). Despite using an especially designed calculator (i.e., Anton) that allows the identification and characterization of the binding pose of a ligand inside a protein,¹⁸ a quantitative description is currently achievable only for fast binders,¹⁹ within the limitation of the force field.

For slower binders (i.e., with a residence time that exceeds the timescale reachable by specific purpose computers, namely, milliseconds) two classes of MD-based approaches can be used. One is based on free-energy perturbation (FEP) approaches.^{20–22} These predict relative differences in binding free energies of ligands for the same target, of the same ligands for a set of different targets, or for mutants of the same target.^{23–27} They correctly take into account entropic effects and they are used widely in drug design campaigns.²⁸ However, likewise molecular docking approaches, they cannot predict (un)binding pathways and become infeasible when significant induced-fit effects take place.^{18,29}

These latter limitations may be overcome by potential mean force calculations. These predict the absolute binding/

unbinding free energy as a function of reaction coordinate(s) or collective variable(s) (CVs) and provide binding/unbinding pathways. Among these methods, metadynamics (MetaD) has been shown to be particularly successful to predict unbinding pathways even in cases where target flexibility plays a significant role.^{30–35} However, in the presence of induced-fit mechanisms, the correct identification of the relevant CVs requires the knowledge of the ligand binding/unbinding processes, in particular, of the conformational rearrangement taking place upon binding. This is not trivial to know a priori when the induced fit is involved. Volume-based MetaD might, in principle, solve this issue using system-independent CVs, allowing the ligand to explore all of the accessible volume surrounding the target.³⁶ However, in practice, the volume needed to exhaustively cover the relevant conformational space with the induced fit (e.g., for including large induced conformational changes upon binding) is very large. As a result, recrossing events (required for the convergence of the free energy) are not very likely, and the calculations are overall not very efficient.

Herein, we have improved volume-based MetaD by: (i) Combining the strengths of system-independent CVs with the benefits of reducing the ligand accessible conformational space to target smaller volumes relevant for the binding. This strategy was already implemented in other MetaD approaches to favor the recrossing events.^{37–40} (ii) Restraining the part of the target that is not directly involved in the binding process. This minimizes the possibility of local unfolding caused by the presence of a localized potential. We show here that this new approach, called localized volume-based metadynamics (LV-MetaD), does not lose the adaptability to conformational changes upon binding, and it dramatically decreases the computational cost needed to obtain convergence. Furthermore, it permits the exploration of possible ligand binding pathways, and it finally overcomes the challenges of ligand-induced fit and slow conformational changes by predicting poses, affinities, and binding mechanisms.

As a test case, we focus on the human MacroD1 and MacroD2 enzymes. These catalyze the hydrolysis of mono-ADP-ribose (ADPr) from protein substrates⁴¹ as well as they efficiently reverse ADPr modification from the 5' or 3' terminal phosphates of DNA and RNA.⁴² ADP-ribosylation is an important post translational modification (PTM) that occurs in multiple key biological processes. In spite of the high pharmacological relevance of these targets,⁴¹ drug design campaigns have been unsatisfying,^{43,44} mostly because they have not been able to identify selective inhibitors for the two enzymes. These would provide important tools to regulate intracellular signaling and regulatory processes, with important applications in neurodegeneration and cancer.⁴⁵

The X-ray structures of the two proteins with and without ADPr (*apo* and *holo* forms, respectively) show that the catalytic sites are deeply buried in the protein.^{46,47} Two active loops (loop1 and loop2, red in Figure 1a) delimit the binding site and function as switch loops to sequester the substrate and they provide sufficient structural flexibility to accommodate diverse types of substrates.⁴⁶ Therefore, not only does ligand binding require substantial changes of the loops' conformations to permit access in the inner cavity to the ligand but also it induces specific loops' conformations upon binding that cannot be predicted *a priori*.⁴⁸

Inspection of the X-ray structures shows that in both enzymes, the ligand ADPr presents an L-shape conformation, and the binding poses of the ligand are the same (Figures 1b and S1). The residues in the two binding sites are also the same on passing from MacroD1 to MacroD2, except that phenylalanine (F272) and valine (V271) in the former are replaced by tyrosine (Y190) and isoleucine (I189) in the latter, respectively (Figure S1). These two different residues in loop2 cause a change in affinity toward ADPr from -9.5 to -8.4 kcal/mol on passing from MacroD2 to MacroD1, respectively. Further mutations of these same two key residues in MacroD2, Y190N and I189R, substantially increase the affinity for ADPr up to -10.1 and -10.3 kcal/mol, respectively.⁴⁸ This experimental evidence suggests a key role of the loops' residues in substrate specificity. However, the molecular mechanisms of such changes in affinity are not known.

Our LV-MetaD scheme turned out to reproduce: (i) The ADPr binding poses of the known complexes in the wild-type (WT) systems and it predicts the unknown ones for Y190N and I189R MacroD2 mutants. (ii) The experimentally measured protein-ADPr binding free energy of the two WT enzymes, along with those of the Y190N and I189R MacroD2

mutants, providing a rationale for the increased affinity of the latter. This information may help rationalize the druggability and substrate specificity of such classes of proteins. Furthermore, our calculations elucidate the molecular interactions induced upon binding at each stage of the binding/unbinding process, paving the way to the design of highly selective inhibitors. Finally, our calculations provide the molecular basis, for the first time, of the impact of all of the mutagenesis experiments on the measured binding affinities so far conducted. Overall, our novel LV-MetaD protocol emerges as a powerful tool to efficiently investigate the induced-fit molecular recognition in ligand/receptor events.

RESULTS AND DISCUSSION

LV-MetaD calculates the ligand substrate (ADPr) binding free energies as a function of apt collective variables. These are the 3D positions of the ligand relative to the binding pocket. The calculations allow predicting substrate ADPr poses and binding pathways in WT MacroD1 and WT, Y190N, and I189R MacroD2. An efficient sampling is achieved by: (i) Limiting the exploration of the ligand within a well-defined volume around the target binding pocket, thus avoiding that most of the sampling time for the ligand is spent in the solvent, a portion of the space not relevant for binding and unbinding. This is achieved by adding a repulsive potential at the boundaries (see the section below). (ii) Adding a constraint on the backbone in parts of the protein not directly involved in the binding/unbinding process (red part in Figure 1c,d). This potential is added because the ligand can be forced to explore volume portions occupied by the host, unfolding the protein.

Choice of the Volume. For WT MacroD2, we set up two different volume shapes to limit the sampling of the ligand. These are either a 1/3 sphere-solid shape (Figure 1c) or a parabolic-solid (Figures 1d and S2) shape. The exploration of the conformational space is satisfactory as several recrossing events occur in both cases (Figure S3). The binding free energies, calculated as specified in the Methods section, turn out to agree well with experimental measurement (Table S1). This confirms the flexibility of the method irrespective of the choice of the volume shape; indeed, the latter can be adapted to maximize sampling efficiency depending on the specific case.

Here, since we observed a slightly better agreement with the experimental data (Table S1) for parabolic-solid volume in MacroD2, we decided to implement this shape for all of the remaining systems. The latter allowed us to compute binding affinities in extremely good agreement with the experimental ones (Table 1).

All of the systems were initially equilibrated by 300 ns of MD, followed by ~ 0.8 μ s-long LV-MetaD simulations. In the next sections, we discuss the impact of induced fit on the binding poses and the (un)binding processes for each system.

WT MacroD2. The calculations on the MacroD2 were based on the *holo* structure of MacroD2 in complex with ADPr.⁵⁰ The free energy shows the presence of four basins (Figures 2 and S4a). The lowest free-energy minima (basins 1 and 2) differ by around 0.5 kcal/mol (i.e., the order of thermal fluctuations, thus they are both populated at room temperature) and represent the bound state. The structure associated with basin 2 reproduces the pose in the *holo* MacroD2 X-ray structure (Figure 3a,b):⁴⁸ the ADPr adenine ring is deeply buried in the highly conserved binding pocket⁴⁸ and forms bifurcated H-bonds with the backbone and the side chain of D78, respectively. The ADPr adenosine ribose forms H-bonds

Table 1. Absolute Free Energy from the Parabolic-Solid LV-MetaD Simulation and Experiment^{46,48,49a}

systems	ΔG_{metaD} (kcal/mol)	ΔG_{corr} (kcal/mol)	ΔG_{std}^0 (kcal/mol)	ΔG_{exp}^0 (kcal/mol)
MacroD2	-7.8 ± 0.3	1.6	-9.4 ± 0.3	-9.5
MacroD1	-6.6 ± 0.3	1.6	-8.2 ± 0.3	-8.4
MacroD2 I189R mutant	-8.1 ± 0.3	1.6	-9.7 ± 0.3	-10.3
MacroD2 Y190N mutant	-8.4 ± 0.3	1.6	-10.0 ± 0.3	-10.1

^a ΔG_{std}^0 is calculated at $T = 298$ K. ΔG_{exp}^0 of WT MacroD2 and its mutant is determined at 298 K, while ΔG_{exp}^0 of WT MacroD1 is detected at 303 K. The ionic strength is 100 mM both in the experimental and computational conditions and the method used for binding assays is isothermal titration calorimetry.

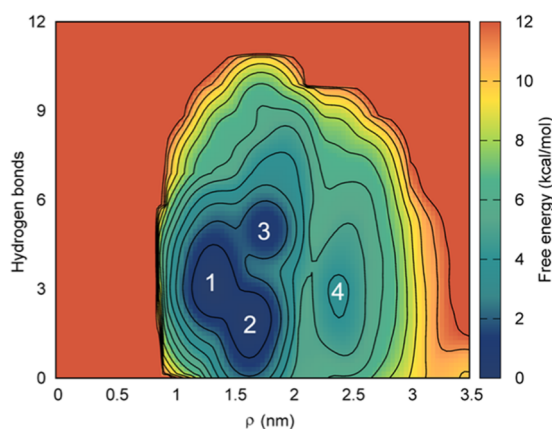


Figure 2. Free-energy surface associated with MacroD2/ADPr binding. This surface is obtained from a reweighting of the deposited bias (originally a 4D surface) to 2 CVs that results more meaningful for the identification of relevant minima, such as ρ (the distance between the center of mass of the target and the center of mass of the ligand) and the number of H-bonds.

with the backbones of C222 and C184. The negatively charged ADPr pyrophosphate moiety located in the cleft of active loop1 and loop2 forms H-bond networks with backbone atoms of I185, S186, T187, G188, and I189 and bifurcated H-bonds with the backbone and the side chain of Y190. The ADPr distal ribose faces the entrance of the binding cavity. It forms H-bonds with the backbone of A90 and the side chain of N92 in loop1.

In basin 1 (Figure 3c), the adenine ring of ADPr no longer interacts with D78 and F224 due to a slight variation of ADPr pose. These interactions are substituted by the formation of hydrophobic contacts with residues V101 and A90 and H-bonds with the V88 backbone (Figure 3c). ADPr adenosine ribose no longer forms hydrogen bonds interactions with the backbone of C222 and C184 but with the backbone of V101. The pyrophosphate moiety forms hydrogen bonds with the side chain of R150 and backbone atoms of residues in loop2 (i.e., G188, I189, and Y190) as well as with G99 (Figure 3c), which is similar to what is observed in basin 2. The distal ribose faces the solvent and forms hydrogen bonds with the backbones of N92 and A93, respectively.

In basin 3 (Figure 3d) ADPr is partially out from the cleft between the two loops and exposed to the solvent. The adenine ring maintains hydrogen bonds with the side chain of D78 and the backbone of I79 and a π -stacking with F224, and

adenosine ribose forms the H-bonds with the C222 and C184 backbones and with the C184 side chain (Figure 3d). A pyrophosphate and a distal ribose moiety rotate above loop2 and are partly sandwiched between helix 9 and helix 10. The pyrophosphate moiety is stabilized by the H-bonds with T187, G188, I189, and N194, while the distal ribose loses all of the interactions with loop1 and loop2. In basin 4, ADPr is located at the entrance of the binding cavity (Figure 3e). The adenine ring is in the cleft formed by loop1 and loop2 and forms π -stacking interactions with Y190 (Figure 3e), and adenosine ribose forms H-bonds with backbones of A91 and A93. The pyrophosphate moiety and distal ribose are completely exposed to the solvent; the former forms H-bonds with the R150 side chain, while the latter forms H-bond interactions with T125, D160, backbone, and side chain, respectively.

From this free-energy landscape, we can gain qualitative insight into the induced-fit ligand (un)binding mechanism: the ADPr adenine ring approaches the binding pocket by interacting with loop1, and then it passes through the cleft between loop1 and loop2 and subsequently reaches the inner binding site in the binding pose (Figure S4b). During the unbinding process, the pyrophosphate moiety and distal ribose of ADPr escape from the groove between loop1 and loop2 and move toward the cleft between helix 9 and helix 10 and then lose all of the interactions with the binding pocket (Figure S4b). In the bound state (basin 2), the loops are as close as 10.8 Å (smallest centroid distance). The transition from the bound to the unbound state, however, triggers a closure and an opening of the loop1–loop2 interface (Figure S4c,d), with their centroid distances ranging from a minimum value of ~ 10.0 Å to a maximum of ~ 14.5 Å (Table S2 and Figure S4c), indicating the presence of ADPr-dependent induced-fit mechanisms. Such movements of the loops cannot be appreciated in the X-ray structure where the loop conformations in *apo* and *holo* states are very similar, with centroid distances of 12.9 and 11.6 Å, respectively (Table S2).⁵⁰ Therefore, our simulations reveal key ligand-dependent events involving loop1 and loop2 impacting on the binding pathway, and, in turn, on the binding affinity (see follow-up paragraphs).

WT MacroD1. The calculations on WT MacroD1 were based on the *apo* X-ray structure instead of the *holo* structure (Figure 4a) to check the robustness of the method in reproducing induced-fit effects of the binding process even starting from a conformation where the ligand is not at its energy minimum. As already discussed above, the LV-MetaD calculations performed on the MacroD1 system turned out to reproduce the free energy of binding experimentally determined (Table 1). The free-energy landscape features only two basins, 1 and 2 (Figures 4b and S5a). Remarkably, basin 1 corresponds to the pose observed in the *holo* MacroD1 X-ray structure (Figure 4c). It features a close conformation as in the X-ray *holo* form, where the centroid distances of loop1 and loop2 are 10.5 and 10.9 Å, respectively (Figure 4a and Table S2),⁴⁶ to be compared with the distance in the X-ray *apo* form where loop1 and loop2 are in an open conformation with a centroid distance of 14.6 Å. This is a remarkable result, as the simulation started from the *apo* form with the ligand docked *a posteriori*, instead of the *holo* form. In basin 1, the adenine ring of ADPr binds to the inner binding pocket establishing π -stacking interaction with F306 and maintains a similar hydrogen-bonding network as in the *holo* MacroD1 structure (Figures 4d and S5c). Adenosine ribose forms H-bonds with the backbone of C266 and the side chain of T269. The

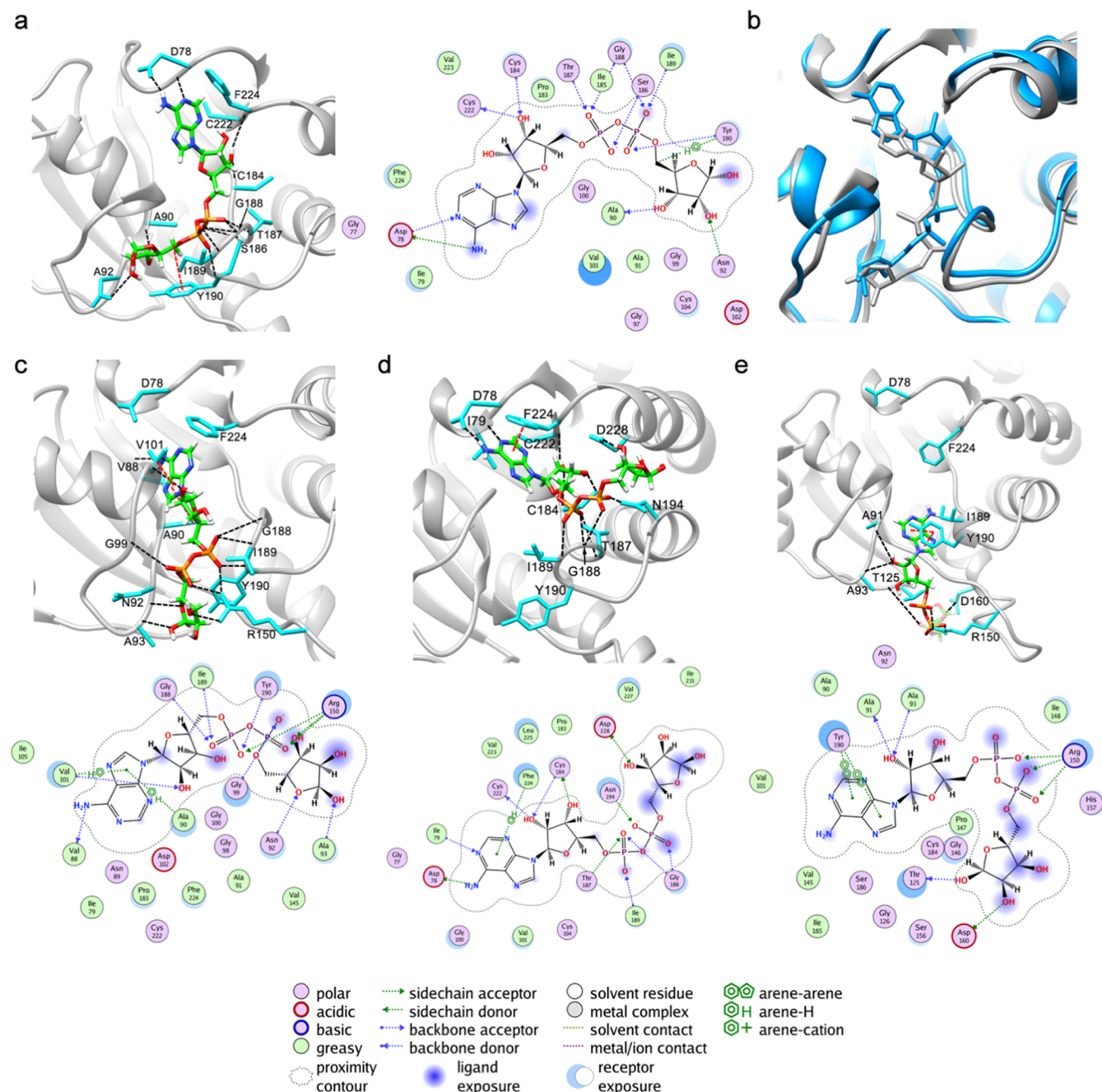


Figure 3. LV-MetaD simulation of ADPr binding to WT MacroD2. (a) Representative conformations of basin 2 (left) and its ligand–protein interaction diagram (right). Bulk water is not shown for clarity hydrogen bonds and hydrophobic interactions are depicted as black and red dashed lines, respectively. (b) Superposition of the *holo* MacroD2 X-ray structure (in gray) with the binding pose of basin 2 (in blue). (c–e) Same as (a) for basins 1, 3, 4. The calculations employ the parabolic-solid scheme shown in Figure 1.

pyrophosphate moiety is stabilized by hydrogen bonds formed with backbones of the residues in the two active loops, i.e., G182, V183, S268, T269, G270, and V271 (Figure 4e). Distal ribose is further away from loop1 and it does not form direct interactions with the two loops. Indeed, it is slightly twisted with respect to the *holo* MacroD1 and MacroD2 X-ray structure (Figure 4c). Basin 2 is ~ 4 kcal/mol higher in free energy with respect to basin 1 (Figure 4b). Here, the *adenine* ring of ADPr is stacked between loop1 and loop2 (Figure S6a). The other parts of the ligand are completely exposed to the solvent above loop1, and they form hydrogen bonds with the

backbone amino groups of G182, G185, and C186, and with the side chain guanidino group of R189 (Figure S6b).

Also in this case, we attempt to gain insight into the (un)binding process by investigating the free-energy landscape. The loop distance change increases relative to the WT MacroD2: the centroid distance ranges between ~ 10.3 and ~ 16.8 Å (Table S2). The adenine ring of ADPr initially approaches the binding site by interacting with helix5 or helix 10, and then the other moieties of the ligand directly lie in the groove between loop1 and loop2. During the unbinding process, the ligand moves its distal ribose and pyrophosphate moieties toward the solvent and then drags the *adenine* moiety

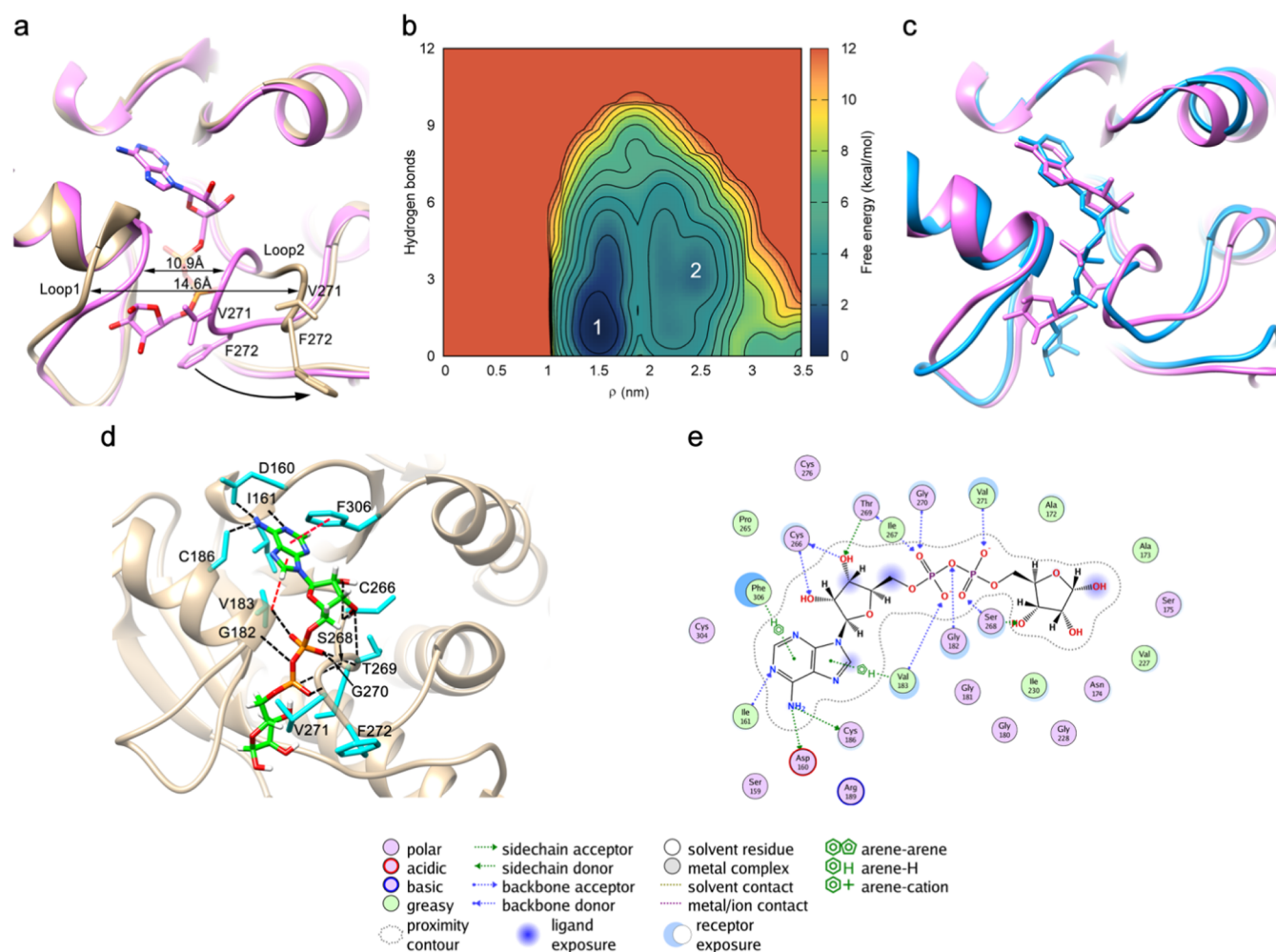


Figure 4. LV-MetaD simulation of ADPr binding to *apo* MacroD1. (a) Superposition of *holo* MacroD1 (PDB code: 6LH4, in purple) and *apo* MacroD1 (PDB code: 2X47, in yellow) X-ray structures. The distance between the center of mass of loop1 and loop2 is highlighted. They consist of 179-GGGGV-183 and 270-GVFG-273 residues, respectively. (b) Free-energy surface as a function of ρ and the number of hydrogen bonds. (c) Superposition of the *holo* MacroD1 X-ray structure (in purple) with the calculated MacroD1-ADPr binding pose of free-energy basin 1 (in blue) highlighted on the free-energy landscape. (d) Representative conformation of free-energy basin 1. Same color code and representation mode of Figure 3 is used. (e) Ligand–protein interaction diagram for the representative structure of free-energy basin 1.

directly from the cleft between the loop1 and loop2 (Figure S5b). This is relevant if we consider that the two residues that discriminate between MacroD2 to MacroD1 (F272 and V271 in MacroD1 are Y190 and I189 in MacroD2) are located in loop2: these differences in the amino acid sequence indeed affect the steric hindrance of the entrance of the binding pocket (V271 in MacroD1 is smaller than I189 in MacroD2), as well as the number of protein–ligand interactions (e.g., the H-bonds between the pyrophosphate moiety and F272 are not present in MacroD1), and, in turn, the ADPr-induced loops' configurations, possibly providing a rationale also for the lower number of intermediate states. Thus, the two residues that are modified on passing from WT MacroD1 to WT MacroD2 cause a loss of interactions between ADPr and loop2, leaving loop1 as the sole player in ligand binding, in contrast to WT MacroD2. Again, these aspects were not identifiable by just the comparison of the *apo* and *holo* crystal structure of MacroD1.

Y190N and I189R MacroD2. No structural information is available for these systems; therefore, they were modeled from WT *holo* MacroD2 (see the Methods section).

Y190N MacroD2. The free-energy profile shows four basins as in the WT protein (Figure 5a). Basin 1 is the lowest

free-energy minimum. The ligand pose is similar to that of WT MacroD2/1 (Figure 5b). The interactions established by the adenine ring and adenosine ribose with the protein are like the ones established in MacroD2 (basin 2). The distal ribose and the pyrophosphate moiety form extensive hydrogen bond networks with residues in the vicinity of loop1 and loop2, i.e., backbones of G100, V101, S186, T187, and G188 and the side chain of D102 (Figure 5c,d). However, the pyrophosphate moiety no longer interacts directly with I189 and N190 as in WT MacroD2. Indeed, we observed that loop2 in Y190N MacroD2 is in an open conformation, different from the close one observed in the WT MacroD2 bound state (Figure 5b). In basin 2, the ligand is located at the entrance of the binding site appearing as in a prebinding state (Figure S7a), while in basin 3, ADPr hangs over the gap between loop1 and loop2 (Figure S7b). In basin 4, ADPr completely escapes from the binding pocket (Figure S7c).

Y190N MacroD2 displays the same number of basins found in WT MacroD2. Interestingly, during the binding/unbinding process, the centroid distances between the two loops sample a broader range of values (from ~ 10.2 to ~ 22.5 Å, Table S2) with respect to WT MacroD2. Notably, the two loops are both

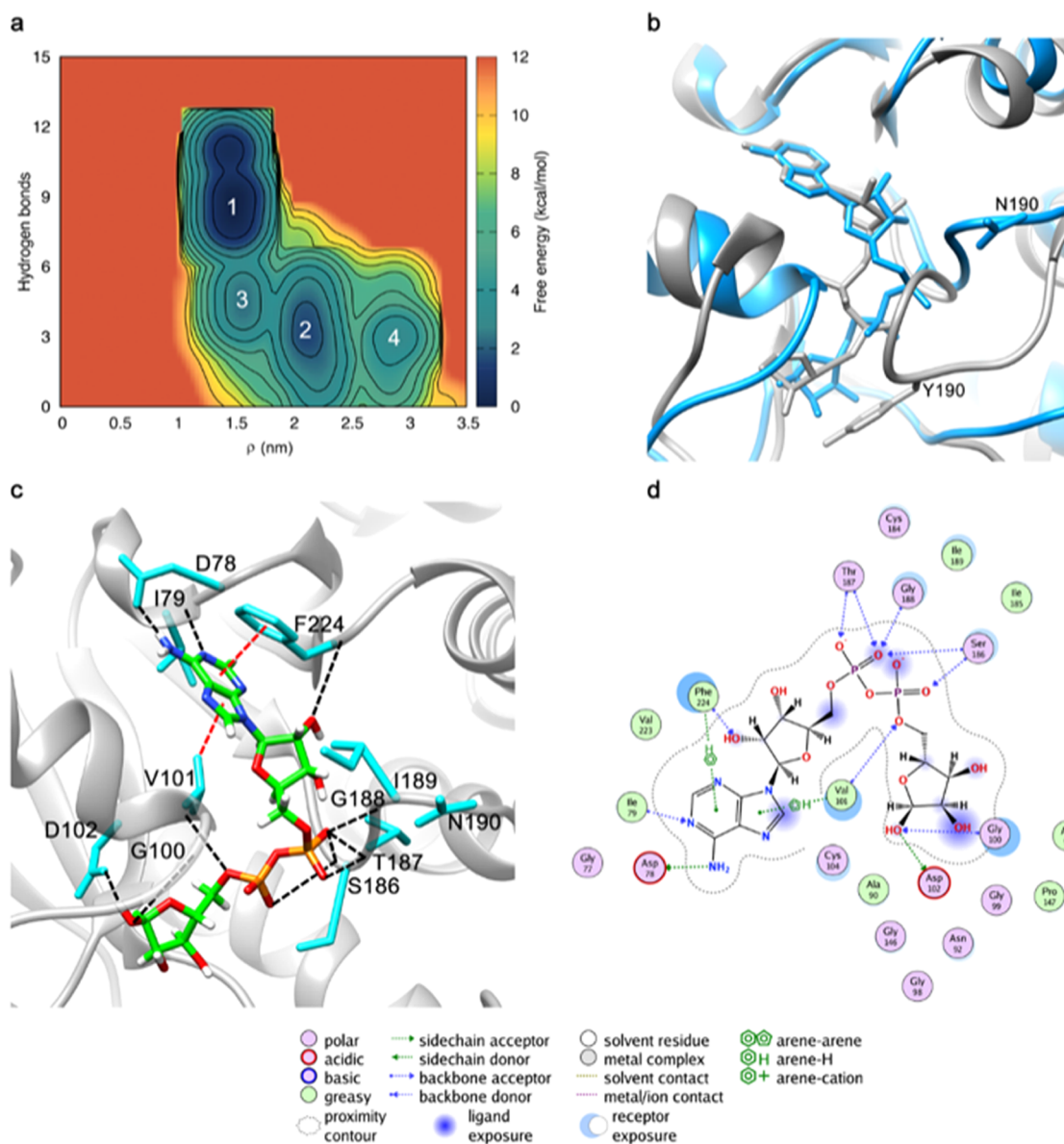


Figure 5. LV-MetaD simulation of ADPr binding to Y190N MacroD2. (a) Free-energy surface as a function of ρ and the number of hydrogen bonds. Here, ρ is the distance of the center of mass between Y190N MacroD2 and ADPr. (b) Superposition of the WT *holo* MacroD2 X-ray structure (in gray) with the binding pose of free-energy basin 1 (in blue) highlighted on the free-energy landscape. (c) Representative conformation of free-energy basin 1 in the binding pocket. The same color code and representation mode of Figure 3 is used. (d) Ligand–protein interaction diagram for the representative structure of free-energy basin 1.

in an open conformation in basins 1–4. Also, ADPr undergoes the same binding process as WT MacroD2 but a different unbinding process. Namely, ADPr escapes from the binding site along the cleft between helix5 and helix6 (Figure S8a,b). These differences on passing from WT MacroD2 to the Y190N mutant might be caused by the smaller side chain in position 190, which from one side increases loop2 flexibility and from the other side decreases the steric hindrance of the cleft between loop1 and loop2. The observed effects could provide a rationale for the increased binding affinity.

I189R MacroD2. Only three basins are observed (Figure 6a), in contrast to four WT MacroD2 and the Y190N mutant. This might be due to the fact that R189 attracts the ligand and locks its orientation inside the binding pocket. Therefore, the conformational freedom of the ligand in the pocket is reduced, possibly lowering the number of intermediate states. The

binding pose of ADPr in basin 1 reproduces the conformation observed in WT MacroD2 and involves almost the same interactions (Figure 6b). The main difference is that the negatively charged *pyrophosphate* moiety is stabilized by a salt bridge interaction with R189 (Figure 6c,d). This new interaction might contribute to the observed higher ADPr binding affinity. Basin 2 corresponds to a predissociated transition state where the ligand attempts to escape from the binding groove into the solvent by moving its distal ribose across loop1. During the motion, the salt bridge interaction with R189 is however maintained (Figure S9a). The binding pose of basin 3 occurs when the ligand leaves basin 2 to reach a more solvent-exposed state (Figures S9b and S10). During the binding/unbinding process, the centroid distance between loop1 and loop2 varies from a maximum distance of ~ 17.5 Å to a minimum distance of ~ 9.8 Å, having 10.6 Å in the bound

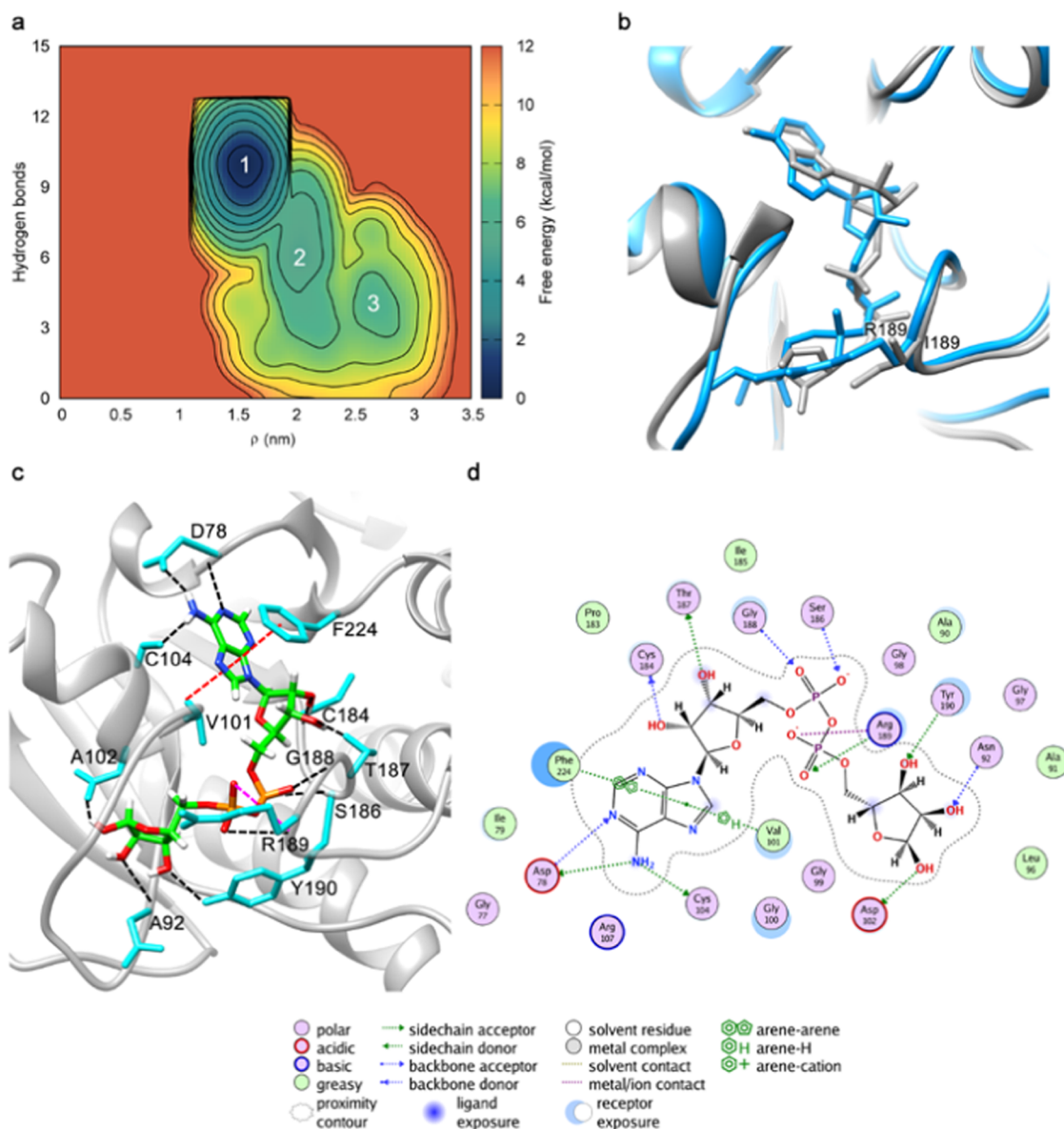


Figure 6. LV-MetaD simulation of ADPr binding to I189R MacroD2. (a) Free-energy surface as a function of ρ and the number of hydrogen bonds. ρ is the distance of the center of mass between the I189R MacroD2 mutant and ADPr. (b) Superposition of the WT MacroD2 crystal structure in complex with ADPr (in gray) with the binding pose of free-energy basin 1 (in blue) highlighted on the free-energy landscape. (c) Representative conformation of basin 1 in the binding pocket. The same color code and representation mode of Figure 3 is used. (d) Ligand–protein interaction diagram for the representative structure of free-energy basin 1.

state (basin 1), comparable to the value observed in the bound states of ADPr in WT MacroD2.

Molecular Basis for the Change in Affinity of MacroD2 Mutants. Based on the affinity of ADPr, the Macro2 mutants have been classified into three groups:⁴⁸ (i) variants featuring low affinity (below 20% of that of the WT), (ii) variants featuring intermediate ADPr affinity (about 40–60% of that of the WT), and (iii) variants featuring WT-like activity and affinity for ADPr or even higher than the WT, i.e., I189R and Y190N, investigated above. Based on our calculations, we attempt here to provide a rationale for the changes in affinity of all of the mutants by visual inspection of the two free-energy minima of WT MacroD2/ADPr (basins 1 and/or 2 in Figure 3), and of the X-ray structures (Figure S1b).

G188E, G100E, and G100E/I189R MacroD2 belong to (i). In basin 2 and in X-ray, the G188 backbone establishes bifurcated H-bonds with the negatively charged pyrophosphate moiety of ADPr; the G100 backbone establishes a single H-bond with the pyrophosphate moiety in X-ray, while in basins 1 and 2, it is in proximity (Figure 3a). Therefore, the G100E and G188E mutations are expected to introduce a bulky negatively charged group near to pyrophosphate, leading to the disruption of the hydrogen-bonding network between the pyrophosphate and the two loops. This may lead to rearrangement of how the ligand is positioned in the catalytic site or it might even prevent binding completely, possibly explaining the dramatic loss of affinity. The affinity of G100E/I189R MacroD2 is larger than that of G188E and G100E MacroD2 but still lower than that of variants in group (ii). The I189R mutation introduces a positively charged group close to

the negatively charged pyrophosphate. Thus, we speculate here that the I189R mutation stabilizes, at least in part, loop2/ligand interactions, leading to a less dramatic change of affinity relative to the WT, as observed experimentally.

D78A, G100S, and N92A/D102A MacroD2 belong to (ii). In basin 2, the D78 side chain and the backbone establish bifurcated H-bonds with ADPr's amino group N6 and N1 atoms (Figure 3a), respectively. The H-bond with N6 is observed also in the X-ray structure (Figure S1). The N92 side chain forms a H-bond with an OH group of distal ribose in both basin 2 and in X-ray (Figures 3a and S1). These interactions are, however, not observed in basin 1 of the same system (where the binding pose of the ligand is still preserved), suggesting that these amino acids are not key for stabilizing ligand binding. D102 in our simulations and in X-ray is in proximity to the distal ribose and the pyrophosphate moiety, respectively, but it does not appear to form key interactions with the ligand. Therefore, it is not possible to understand whether the D102A mutant might have an effect if uncoupled from N92A. G100, as said above, only establishes a single H-bond with the pyrophosphate moiety in the X-ray that is not observed in basins 1 and 2. Indeed, G100S has only a mild effect on the binding affinity. The smaller impact of G100S with respect to G100E described above can be rationalized by the smaller steric hindrance and missing net charge of serine with respect to glutamate.

I189R, Y190N, F224A, and G99E MacroD2 belong to (iii). The rationale for the increased affinity of I189R and Y190N has been given above. F224A and G99E MacroD2 feature affinities close to that of the WT. Notably, while in X-ray, these residues feature a π -stacking with the adenine ring and H-bond with distal ribose, respectively. These interactions are not observed in basins 2 and 1 of our simulations. G99 is indeed solvent-exposed in our simulations and therefore its substitution for E is not expected to affect ligand binding, as confirmed by the experiments (Figure 3a). F224 is only in proximity to the ligand, therefore its substitution with A is not expected to play a key role for the affinity, consistently with experimental evidence. Flexibility analysis of the ligand in the simulated systems is shown in Figure S11.

Overall, our simulations can provide a rationale for the different affinities observed in mutagenesis experiments, not fully emerging just from the visual inspection of the X-ray structure.

CONCLUSIONS

To summarize, we have presented a new method, called localized volume-based MetaD scheme (LV-MetaD), to predict poses, affinities, and (un)binding mechanisms of small ligands targeting proteins, even in cases where induced-fit effects are significant. The method exploits the advantages of system-independent CVs combined with a restricted conformational space sampling. To the best of our knowledge, this is the first time that such a general scheme has been developed. LV-MetaD has been applied to substrate (ADPr) binding to macrodomain enzymes, WT MacroD1, and WT MacroD2 and their binding to ADPr, for which we know that induced-fit effects are present: two highly flexible loops regulate the access to the active site and rearrange in a specific manner depending on the ligand.⁴⁶ Simulations have also been extended to Y190N MacroD2 and I189R MacroD2 mutants, which are the only mutants for which the affinity increases.

The calculated absolute binding free-energy differences of the ligand binding to the WT proteins turn out to compare well with experiments. The scheme also reproduces the increased affinity of the mutants with respect to WT MacroD2, although the difference in the affinity between Y190N MacroD2 and I189R MacroD2 is beyond the precision of the method. Taken together, these results establish the accuracy of the calculations.

The calculations further predict the ligand poses of ADPr in MacroD2 mutants, without prior knowledge of it and depict the differences in the binding/unbinding pathways with respect to WT MacroD2. In addition, they provide a rationale for the impact of the other mutations for their consequences of affinity. Overall, our simulations confirm that the conformational rearrangements of the binding pocket are induced by ADPr binding and they demonstrate the capability of the LV-MetaD protocol, given the possible limitations of classical force fields and/or within the statistical uncertainties of the experimental data, to accurately predict the binding free energy, the binding conformation, and characterize the induced-fit binding mechanisms. Moreover, no prior knowledge of the binding pose is required.

The scheme is entirely general, and it could be applied to investigate ligand binding to other enzymes and receptors for which induced fit is important. The free energy is described by a set of collective variables that can be set up according to the shape of the binding pocket. The application of this scheme with a new and efficient form of CV-based enhanced sampling protocols, such as the on-the-fly probability enhanced sampling (OPES),⁵¹ combined with the new computational capabilities given by the exascale computing project, can finally pave the way for a totally general *in silico* drug discovery approach based on molecular simulations.

METHODS

System Preparation. The crystal structure of human mono-ADPr hydrolase macrodomains has been reported in the literature: the first is MacroD2 in complex with ADPr (PDB access code 4IQY), resolved at a 1.5 Å atomic resolution.⁴⁸ The crystal structure has two protein molecules, by aligning them, the RMSD is less than 0.1 Å, revealing it has two nearly identical structures. Hence, to reduce the computational cost, we performed all of the simulations using the monomer molecule. The second is the MacroD1 monomer in the apo state at a 1.7 Å resolution (PDB access code 2X47)⁴⁷ and MacroD1_ADPr complex at a 2.0 Å resolution (PDB access code 6LH4).⁴⁶ The two monomers involve a general macrodomain fold as a three-layered α - β - α sandwich structure with a central six-stranded β sheet and are evolutionarily conserved protein with an almost identical binding site. To test the capability of the LV-MetaD scheme to predict the correct binding pose in the cases where only the apo structure of the target is available, the initial structure of holo MacroD1 was built by docking ADPr to the available apo MacroD1 X-ray structure⁴⁷ instead of using the holo MacroD1 X-ray structure.⁴⁶ Prior to docking the ligand to the active site, the ligand's geometry was optimized using the LigPrep module; the protein was processed using the Protein Preparation Wizard. The docking studies were carried out using Glide software (Maestro 12.0, Schrödinger, LLC). The default setup was used for all of these steps. Additional two mutants of MacroD2 were obtained by replacing I189 with arginine and Y190 with asparagine, respectively, using UCSF Chimera.⁵²

The missing loops in these two protein crystallographic structures are far away from the binding site and they were reconstructed using the Protein Preparation Wizard of Schrödinger.⁵³ Due to the lack of the structure of the MacroD2 mutant bound to ADPr, the structure of the WT MacroD2 complex was employed as a reference model for the MacroD2 mutant binding mode.

Each protein-ADPr complex system was performed with AMBER ff99SB-ILDN force field for the protein and ions,⁵⁴ and was solvated with TIP3P explicit water molecules in a periodic cubic box, in which the protein surface was 20 Å far away from the periodic box edge.⁵⁵ An additional appropriate number of sodium and chloride counterions were added to neutralize the charge of the whole system and to achieve a physiological salt concentration of ~100 mM, which was consistent with the experimental ionic force used to determine the binding constant.^{48,49} Topology for the ligand was constructed using the general AMBER force field (GAFF)⁵⁶ and its partial atomic charges were parameterized with the AM1-BCC semiempirical method.⁵⁷

MD Simulations. All of the simulations were performed using GROMACS 2019.2⁵⁸ patched with the PLUMED 2.6.^{59,60} Long-range electrostatic interactions were calculated with the particle mesh Ewald (PME) scheme using a grid spacing of 1.2 Å and the distance cutoff for the short-range electrostatic interactions and van der Waals was set to 12 Å.⁶¹ All of the bonds including the hydrogen bonds were restrained with the LINCS algorithm.⁶² The velocity rescale thermostat⁶³ with the solvent and the solute coupled to separate heat baths was employed for the temperature regulation and the pressure was controlled using a Parrinello–Rahman barostat.⁶⁴ After an energy minimization of the solvent with the steepest descent algorithm, the temperature of each system was gradually increased from 0 to 298 K in 1 ns of MD. Keeping the position restraints for the protein and ligand, the system was equilibrated at 298 K for 10 ns in the NVT ensemble; after this step, we released the ligand, and the system was equilibrated by 10 ns in the NPT ensemble. Finally, a long equilibration of 300 ns without any restraint was performed before the production MetaD simulation.

Metadynamics Simulations. Here, we developed the localized volume-based metadynamics scheme on the base of funnel-shaped MetaD and spherical volume-based MetaD. We chose two different restraining volume shapes. The first restraining applied was 1/3 sphere-shaped. As in the original volume-based protocol, we biased the spherical coordinate system: CV1- ρ , the distance between the center of mass of the ligand and protein, CV2- φ , the polar angle measured from the z-axis, and CV3- θ , the azimuthal angle of its orthogonal projection on the x - y plane^{33,65} (Figure 1c). To find an appropriate size of the restraining potential (i.e., that contains a reasonably solvated set of conformation for the ligand), we first performed a calculation to estimate the distance between the protein and ligand to avoid any host–guest interactions. After this estimate, the value of ρ was restricted to 35 Å and the value of φ was limited to $\pi/3$. The second restraining potential was shaped as a rotational parabolic-solid. Like in the first case, we biased three CVs, ρ , θ , and σ , where ρ and θ was the same spherical coordinate as mentioned above, and σ was a series of confocal parabolas centered on the center of mass of protein-heavy atoms (Figure 1d). The value of σ ($\sigma = 1$) depends on the restraining space, which should totally include the protein

pocket and allow the ligand flexible exploration of the binding/unbinding orientation.

As previously said, a repulsive potential was applied at the edge of the designed volume, impeding the ligand from visiting less relevant regions in the solvated state. In our case, the restraining potential completely enveloped the binding pocket and was applied along the direction defined from the centers of mass of the whole protein and the binding pocket alone (aligned to the z-axis in the initial structure). We also applied a restraining bias to limit the translation of the center of mass of the protein (restrained a radius of 10 Å). We finally applied a restraining potential on the backbone of the residues that are not involved in the protein–ligand binding process (RMSD < 1 Å).

$$V_{\text{RMSD}}(t) = \begin{cases} 0 & \text{if RMSD}(t) \leq 1 \text{ \AA} \\ k \cdot \text{RMSD}(t)^2 & \text{if RMSD}(t) > 1 \text{ \AA} \end{cases} \quad (1)$$

where k is the harmonic constant, equal to 41.5 (kcal/mol)/Å². To include the atoms that can preserve the system from unfolding events without hampering possible rearrangements in the binding site proximity, we calculated the RMSD considering the residues that are farther than 6 Å from the binding site.

In our case, the MetaD simulations with these two LV-MetaD schemes in the WT MacroD2 systems led to very similar results, which further verified the reliability of both the shapes (Table S1). Given that ADPr binds to a buried and open active site, the MacroD1 and two MacroD2 variant systems were performed with parabolic-solid-shaped potentials. In the well-tempered MetaD simulation, Gaussian hills with a height of 1.2 kJ/mol were deposited every ps, and the Gaussian functions were rescaled using a bias factor of 20, while the widths of Gaussians were set as 1.0 Å, $\pi/8$ rad, and 0.04 for the three CVs— ρ , θ , and σ , respectively. For all of the systems, 0.8 μ s MetaD simulations were conducted and their convergence were achieved approximately at 0.6 μ s (Figure S3a). All of the errors for free-energy differences were evaluated using a block average analysis. The errors were finally determined with a time interval of 50 ns, where they reached a plateau and no longer changed with the block length ranging from 5 to 100 ns.

The representative structure corresponding to the lowest energy basin of each system was extracted along the free-energy surface using the Metadynminer3d.⁶⁶ It has been equilibrated by unbiased MD of 15 ns simulation time. The flexibility of the binding site and the ligand atoms was evaluated and is reported in Figure S11.

Free-Energy Calculation. The experimental absolute free-energy values were calculated from the equilibrium dissociation constant K_d through the formula³⁵

$$\Delta G_{\text{exp}}^0 = -RT \cdot \ln \left(\frac{K_d}{[L]^0} \right) \quad (2)$$

where R is the gas constant, T is the system temperature, $[L]^0$ is the standard state concentration of 1 mol/L, and K_d is the ligand-binding affinity. In the MetaD simulation, the binding free energy (ΔG_{metaD}) was the free-energy difference between the bound and unbound states.

However, the application of restraining potential caused a loss of the translational degrees of freedom of the ligand in the solvated state, which would need to be corrected when calculating the absolute free energy (ΔG_{std}^0).^{2,36,37,67}

$$\Delta G^0 = \Delta G_{\text{metaD}} - RT \cdot \log\left(\frac{L}{L^0}\right)$$

$$= (G_{\text{bound}} - G_{\text{unbound}}) - RT \cdot \log\left(\frac{1/N_A}{V_{\text{space}} - V_{\text{prot}}}\right) \quad (3)$$

where R is the gas constant, T is the system temperature, $[L]$ is the concentration of the ligand in the restraining space, $N_A = 6.022 \times 10^{23}$ is the Avogadro constant, V_{space} is the volume of the restraining potential, and V_{prot} is the volume of protein inside the restraint. The bound state was identified as the free-energy minimum of the landscape and the unbound state as the area of the free-energy surface with a distance between the protein center of mass and the ligand larger than 30 Å. Further information on the calculation of the volume of a parabolic-solid can be found in Figure S1.

■ ASSOCIATED CONTENT

SI Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.jctc.1c00649>.

Details on the protein and system model, details on the computational protocol, binding free energy data, additional data of four modeling systems, Tables S1 and S2, and Figures S1–S11 (PDF)

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Notes

The authors declare no competing financial interest. All of the data and PLUMED input files required to reproduce the results reported in this paper are available on PLUMED-NEST (www.plumed-nest.org), the public repository of the PLUMED consortium⁶⁸ as plumID: 21.018.

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