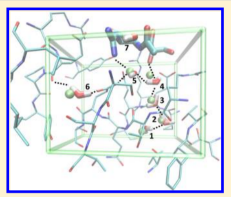
Strategies to Calculate Water Binding Free Energies in Protein-Ligand Complexes

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Supporting Information

ABSTRACT: Water molecules are commonplace in protein binding pockets, where they can typically form a complex between the protein and a ligand or become displaced upon ligand binding. As a result, it is often of great interest to establish both the binding free energy and location of such molecules. Several approaches to predicting the location and affinity of water molecules to proteins have been proposed and utilized in the literature, although it is often unclear which method should be used under what circumstances. We report here a comparison between three such methodologies, Just Add Water Molecules (JAWS), Grand Canonical Monte Carlo (GCMC), and double-decoupling, in the hope of understanding the advantages and limitations of each method when applied to enclosed binding sites. As a result, we have adapted the JAWS scoring procedure, allowing the binding free energies of strongly bound water molecules to be calculated to a high degree of accuracy, requiring significantly less computational effort than more rigorous



approaches. The combination of JAWS and GCMC offers a route to a rapid scheme capable of both locating and scoring water molecules for rational drug design.

INTRODUCTION

The role of water molecules in protein-ligand structures and drug design has become of considerable interest in recent years. 1-5 Fuelled by the early work of Poornima and Dean in 1995, 6-8 water molecules are increasingly being included in a number of stages in the drug discovery process, primarily in virtual screening and docking approaches. 9–11 Traditionally, it has been thought that there are two major roles that water molecules play in ligand binding. The first is to stabilize a protein-ligand complex through creating a hydrogen bonding network, as seen in the binding of zanamivir to N9-neuraminidase. 12,13 The second is the ability of a water molecule to be displaced upon ligand binding, demonstrated through the development of cyclic urea analogues to target HIV-1 protease. ¹⁴ This is typically advantageous since the release of an ordered water molecule into the bulk carries an entropic gain, coupled with an enthalpic gain of strong protein-ligand interactions. 15

More recent studies have helped to shed even more light upon the multiple, and often complex, roles which water molecules play in protein binding sites. Seemingly subtle changes in water-based hydrogen bonding networks have been shown to affect ligand-protein interaction energies, 16 highlighting the need for an accurate representation of solvation within a protein binding site. Studies by Setny, Baron, and McCammon 17,18 have shown that water molecules play an active role in determining ligand-protein binding or rejection, whether this is due to mediating direct interactions or providing an electrostatic screening effect. A recent communication by

Shan et al. has also shown that water molecules can form a solvation shell between the protein and ligand in the binding site prior to the binding event, with the formation of this solvation shell providing a kinetic barrier to binding. 19 Bren and Janežič have developed a molecular dynamics method which decouples the individual degrees of freedom of water molecules, allowing for the simulation of waters at any given combination of rotational, translational and vibrational temperatures.²⁰ Such an approach allows for the identification of the delicate interactions between water molecules and their environment and has shown promise in the calculation of the hydration free energy of water and predicting the structural properties of water under different excited states and environments.

While individual, and clusters of, water molecules can directly influence ligand binding events, the structure of the water networks can also play a critical role. Experimental studies by Homans²¹ have challenged the commonly held belief that protein-binding sites are fully solvated. Applying isothermal titration calorimetry to the major urinary protein system alongside molecular dynamics (MD), it was found that the pocket was suboptimally hydrated, with ligand binding driven by favorable protein-solute dispersion interactions rather than the expected entropic gain in displacing water from the pocket. Upon the basis of this, it is argued that shape complementarity between ligand and protein is a viable method for drug design

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to target the suboptimal hydration patterns. A similar argument has been proposed by Englert et al. looking at the binding of phosphonamidate to thermolysin.²²

Owing to the complex roles which water can perform in protein binding sites, 23 the reliable incorporation of water molecules into computational drug design is of critical importance. Indeed, studies have shown that incorporating water molecules into docking and virtual screening approaches can dramatically improve the predictions formed when analyzing the predicted poses. 10,11 Crystallographic approaches have long been used to identify waters in and around protein binding sites, although this approach is often limited. Carugo has suggested that protein resolution is typically a limiting factor in determining the number of water molecules in a protein structure.²⁴ Through analysis of 873 known crystal structures, the authors indicated that, on average, a protein structure with a resolution of 2 Å had one water molecule per residue, while at a resolution of 1.0 Å around 1.6-1.7 waters are resolved. The same behavior has been noted by Abel et al.²⁵ Another issue with relying on crystallographic methods to predict waters lies in the role of the crystallographer. It has been demonstrated that two independently resolved structures of the transforming growth factor- β 2 were found to have a different number of crystallographic waters with varying temperature factors²⁶ when analyzed by different crystallographers, suggesting that the addition of water molecules into a crystal structure can be problematic.

It is therefore apparent that relying upon crystallographic evidence is not always sufficient if the location of water molecules in protein binding sites is of interest. As such, various simulation methods have been developed to locate water molecules in protein binding sites. The Grand Canonical Monte Carlo method (GCMC)^{27,28} is capable of locating water molecules in protein binding sites using the μVT ensemble.^{29,30} During a GCMC simulation, the number of water molecules is allowed to fluctuate according to the defined chemical potential μ . The chemical potential can be related directly to the binding free energy of the molecule, allowing both the location and affinity of water molecules to be found during a simulation.³¹ One major drawback to the method lies in the poor acceptance rates, with water molecule insertions typically accepted with a probability of <1% even if cavity-biasing 32,33 and configurational bias 32 schemes are utilized.

Based upon inhomogeneous fluid solvation theory (IFST), ^{35–38} the WaterMap method has shown promise in both locating water molecules in protein binding sites and also assessing their free energies. ^{39,40} An MD simulation is performed which tracks the location of water molecules, followed by a clustering procedure which places water molecules. These sites are then subjected to the IFST analysis to estimate the enthalpy and entropy of each water site. One problem with this approach is that low density solvent regions are not accounted for and that the clustering does not discriminate between short and long-lived high-density hydration sites. In addition, sampling occluded binding sites can be problematic, ⁴¹ due to the long time scales required to allow the passage of water between the bulk and the binding site. The WaterMap methodology has been used on a number of systems, including a range of kinases, ⁴² the PDZ domain, ⁴³ and factor Xa. ⁴⁰

To help sample solvent-inaccessible protein binding sites, the Just Add Water Molecules (JAWS) method was developed. The JAWS method is based upon λ -dynamics and is capable of both locating the position of waters within a protein binding site and also providing an estimate for the binding affinity of

these waters compared to the bulk. The approach works by initially simulating so-called θ water molecules which can appear and disappear across a user-defined volume typically centered on the binding site, a process referred to as JAWS stage 1. θ is an energy scaling parameter which controls the interaction energy $U_i(r)$ between θ -water i and the rest of the system. If the value of θ_i is 0 then the molecule acts like a ghost particle and does not interact with the system. Equally, if $\theta_i=1$ then the molecule interacts fully with its surroundings. A simulation is performed whereby the set of θ parameters are sampled using a Metropolis Monte Carlo scheme, alongside full motion of the protein side chains and any adjacent ligands, which results in a population density map of favorable hydration sites on the grid.

The resulting population densities are then clustered into an integer number of hydration sites, with water molecules constrained at each site using a hardwall potential. A biasing potential based upon the hydration free energy of water, together with a correction term for the incorporation of the constraint, is applied to each molecule, and a new simulation performed to observe the probability of a water molecule experiencing states with high ($\theta > 0.95$) or low ($\theta < 0.05$) values, a process termed JAWS stage 2. The cubic volume of the constraint is typically close to that of the volume which a water molecule occupies in the bulk, with the hardwall potential necessary to prevent the water molecule from leaving its site as the intermolecular interactions are removed. The ratio of these probabilities is then used to derive the binding free energy. As with the WaterMap approach, one drawback lies in the clustering of population density, while another lies in the adequate sampling of the θ < 0.05 state for strongly bound water molecules, preventing an estimate of the binding free energy.⁴⁵ The JAWS methodology has been used by Michel et al. to evaluate the energetics of water displacement from three protein binding sites, finding that ligand affinity changes strongly correlate with the binding free energy of the displaced water molecules. 46 Luccarelli et al. have employed the JAWS methodology to optimize the placement of water molecules in the binding site of p38a MAP kinase. They reported significant improvements in the accuracy of subsequent MC/FEP relative binding free energy predictions for 17 ligands in comparison with standard solvent equilibration protocols, highlighting the importance of accurate water placement for structure-based molecular modeling.4

Although the method is not capable of predicting the location of water molecules, the double-decoupling method⁴⁸ is seen as the "gold standard" in calculating the binding free energy of water molecules in protein—ligand complexes, owing to its rigorous methodology. The method involves running two simulations, whereby a water molecule is first decoupled from a box of bulk water, with a second simulation decoupling the water molecule from the protein—ligand complex. The free energy changes from these simulations can then be used to construct a free energy cycle, from which the binding free energy can be found.

Barillari et al. have utilized double-decoupling to help understand the nature of water molecules which can be displaced on protein—ligand binding and those which cannot. 49 Utilizing the method with Replica Exchange Thermodynamic Integration (RETI), 50,51 the study focused on understanding whether the binding affinity of a water molecule was related to its propensity to be displaced. The paper demonstrated that, on average, water molecules which are more tightly bound are less likely to

be displaced than those which are more weakly bound. With this knowledge the medicinal chemist can decide whether to target a particular water molecule for displacement or to try to design a ligand which is capable of utilizing the hydrogen bonding opportunities afforded by that water. The double-decoupling method has also been used more recently by Fadda to investigate the stability of conserved water molecules in Concanavalin A.⁴

Given the different approaches reported in the literature, it can be difficult to determine which method should be used given a particular problem. In addition, no study has ever addressed whether the methods themselves give comparable results. For example, will the binding free energy of a water molecule calculated by GCMC be comparable to that calculated by double-decoupling? In this study three freely available methods are compared, GCMC, JAWS, and double-decoupling, and applied to N9 neuraminidase. This allows the various methods to be fairly assessed and highlights the advantages and disadvantages of each method for locating and scoring water molecules. In the case of JAWS, a solution to the problem of evaluating the binding free energies of tightly bound water molecules is proposed.

METHODS

System Setup. The crystal structure chosen for the simulations was 1*nnc* (resolution = 1.80 Å).¹² Polar hydrogens were added onto the structure using the HBONDS option in whatif v7.0 (the protonation states of the ionizable residues can be found in the Supporting Information),⁵² with nonpolar hydrogens added using LEaP. The zanamivir ligand was parametrized using the antechamber module in AMBER, with the partial charges assigned using the AM1-BCC model.⁵³ The partial charges obtained (found in the Supporting Information) are broadly similar to those reported in a molecular dynamics study by Udommaneethanakit et al.,⁵⁴ which used the RESP approach to assign charges. Any differences are due to the different charge-fitting procedure used in the two studies, making direct comparisons difficult.

The complex was then minimized using the Sander module of AMBER to remove bad contacts, using an igb keyword of 1. To reduce the computational cost, only protein residues that have a heavy atom within 15 Å of zanamivir were retained. Crystallographic waters within this region were retained, except for those which were predicted in this study. The complex was solvated by a sphere of TIP4P water molecules of 23 Å radius centered upon zanamivir. To prevent evaporation, a half-harmonic potential with a 1.5 kcal mol⁻¹ force constant was applied to all water molecules whose oxygen atom distance to the crystallographic zanamivir center of geometry was greater than 23 Å. The resulting complex was then equilibrated for 10 million moves in the *NVT* ensemble to remove bad contacts. During the equilibration, solvent moves were attempted with a probability of 85.7%, protein side-chain moves with a probability of 12.9%, and solute moves with a probability of 1.4%. For the free energy calculations, the amber99 force field was used, 56 with a temperature of 25 °C and a residue based nonbonded cutoff of 10 Å, feathered over the last 0.5 Å. Brunsteiner has demonstrated that similar hydration free energies of water are obtained for group-based cutoff schemes as Ewald-summation schemes, 57 and consequently long-range electrostatic interactions were not modeled in this work. Any ligands used in the studies were modeled using the GAFF force field.58

All simulations were performed using a modified version of the Monte Carlo code, ProtoMS 2.2. For the free energy calculations, the protein backbone was constrained while only the side chains were sampled. Constraining the backbone allows for consistency when comparing the different free energy methods, and the strong interactions between zanamivir, neuraminidase, and the bound waters in the pocket suggest that the inclusion of backbone motion in these simulations is unlikely to induce a statistically significant shift in the reported water binding free energies. For more flexible and open binding pockets, the inclusion of backbone motion is necessary to calculate reliable binding free energies.

JAWS Protocol. The JAWS stage 1 simulation was performed upon the entire binding site, encompassing a grid of size 13 Å \times 9 Å \times 9 Å. TIP4P JAWS waters (48) were added to the simulation region, 55 with these molecules allowed to move freely around the grid region for one million moves while turned off ($\theta = 0$). Unless stated otherwise, the θ threshold applied for water molecules being classed as "on" was 0.95 and "off" as 0.05. Statistics were then collected on the grid region for 40 million MC moves using a grid spacing of 1 Å, in line with the original JAWS study.⁴¹ The resulting data were analyzed using AstexViewer,⁶⁰ and each grid point normalized according to the number density of the most populated grid coordinate. During the simulation, the JAWS waters were allowed to move and sample their associated θ_i value, with full sampling of the ligand angles and dihedrals and bulk solvent performed. The bond angles and torsions for the side chains of residues within 10 Å of any heavy atom of zanamivir were also sampled, with the protein backbone constrained throughout the simulation. For the JAWS stage 1 simulations, solvent moves were attempted with a probability of 23%, protein side-chain moves with a probability of 3.6%, and solute moves with a probability of 0.4%. Variations in θ_i were attempted with a probability of 50%, in line with the original JAWS study, 41 with translations and rotations of the JAWS waters attempted with a probability of 23%.

The JAWS stage 1 simulation identified seven hydration sites which were then used as starting points for the free energy methods.

JAWS stage 2 simulations were performed by placing a 3 Å \times 3 Å \times 3 Å grid over the water molecule of interest. The biasing potential, as described in eqs 1 and 2, was added for the θ -water to the potential energy, and statistics regarding the value of θ collected for 40 million MC moves.

$$V(\theta_i) = (-\Delta G_{\text{hyd}} + \Delta G_{\text{constr}}(\text{ideal, site } i))\theta_i$$
 (1)

In eq 1, $\Delta G_{\rm constr}$ is the free energy for constraining an ideal particle in a volume of $V^{\rm constr}$ instead of the bulk, $V^{\rm c.48}$

$$\Delta G_{\rm constr} = -k_{\rm B}T \ln \frac{V^{\rm constr}}{V^{\rm o}} \tag{2}$$

The hydration free energy of water used in the biasing potential, $\Delta G_{\rm hyd}$, was taken to be $-6.4~{\rm kcal~mol}^{-1}$ in line with previous studies. A binding free energy for the water molecule was found from the ratio of probabilities of observing a θ -water at high (θ > 0.95) and low (θ < 0.05) θ values, using eq 3.

$$\Delta G_{\rm bind}({\rm water, \ site \ }i) = -k_{\rm B}T \, \ln\!\left(\frac{P(\theta_i \to 1)}{P(\theta_i \to 0)}\right)$$
 (3)

In eq 3, $k_{\rm B}$ is the Boltzmann constant and T is the temperature of the simulation. The θ thresholds for the high and low θ

states are arbitrary and consistent with the original JAWS study.⁴¹ The dependence of the calculated free energies on the choice of threshold is investigated later in this paper.

For the JAWS stage 2 simulations, solvent moves were attempted with a probability of 23%, protein side-chain moves with a probability of 3.6%, and solute moves with a probability of 0.4%. Variations in θ_i were attempted with a probability of 50%, consistent with the original JAWS study,⁴¹ with translations and rotations of the isolated JAWS water attempted with a probability of 23%.

Double-Decoupling Protocol. Double-decoupling⁴⁸ simulations were performed using Replica Exchange Thermodynamic Integration (RETI)^{50,51} and the coordinates found from the JAWS stage 1 simulation. The binding free energy of a water molecule was found in two stages using a single topology approach: first the electrostatic terms between the water molecule and its environment were linearly perturbed to zero, followed by a gradual linear reduction in the Lennard-Jones parameters on the oxygen atom to reduce its size to zero. The water molecules were constrained by a hardwall constraint of radius 1.8 Å to allow direct comparison with the JAWS hardwall which has a similar size. The hardwall was applied to only the water in question and forbids it from leaving this spherical region. Furthermore, other water molecules, solute atoms, and protein atoms were not permitted to diffuse into this excluded region. As shown in eq 4, the volume of this spherical hardwall, Veff, can be calculated to be 24.43 Å³, which is of similar size to the cubic 27 Å³ hardwall used in JAWS stage 2

$$\Delta G_{\text{rest}} = -k_{\text{B}} T \ln \frac{V^{\text{eff}}}{V^{\text{o}}}$$
(4)

In eq 4, $k_{\rm B}$ is the Boltzmann constant, T is the temperature of the simulation, $V^{\rm eff}$ is the volume occupied by the hardwall, and V^0 is the standard state volume of water, 29.89 ų at 55.56 M. From this the free energy correction term of the hardwall, $\Delta G_{\rm rest}$, for double decoupling simulations can be found to be +0.12 kcal mol⁻¹.

For both the electrostatic and Lennard-Jones decoupling simulations, 16 equally spaced λ windows were used with a value of $\Delta\lambda$ of 0.001 in conjunction with RETI. The decoupling of both the electrostatic and Lennard-Jones interactions was performed in 40 million MC steps divided into 400 blocks of 100 K steps each. Data were collected and averaged over the last 30 million steps for both sets of simulations. At the end of the simulation, the computed free energies for the decoupling of the electrostatic terms of the molecule and decoupling the Lennard-Jones terms were summed, to give a value for ΔG_{comp} .

Having calculated the values of ΔG_{comp} and ΔG_{rest} the binding free energy of a water molecule, ΔG_{abs} , was found using eqs 5 and 6.

$$\begin{split} S_{\rm sol} &\to S_{\rm gas} & -\Delta G_{\rm hyd} \\ RS_{\rm sol} &\to S_{\rm gas} + R_{\rm sol} & \Delta G_{\rm dec} \\ R_{\rm sol} &+ S_{\rm sol} &\to RS_{\rm sol} & \Delta G_{\rm abs} = -\Delta G_{\rm hyd} - \Delta G_{\rm dec} \end{split} \tag{5}$$

$$\Delta G_{\text{dec}} = \Delta G_{\text{comp}} + \Delta G_{\text{rest}} - k_{\text{B}} T \ln \frac{\sigma_{\text{RS}}}{\sigma_{\text{R}} \sigma_{\text{S}}} + P^{0} (V_{R} - V_{\text{RS}})$$
(6)

The third term in eq 6 is a symmetry related term. T is the temperature, σ_{RS} is the symmetry number of the complex, σ_{R} is

the symmetry number of the protein, and σ_S is the symmetry number of water. Water has a symmetry number of 2, and since the other two terms have a symmetry of 1, the term can be found to be -0.4 kcal mol⁻¹. The final term in eq 6 is taken to be negligible under standard pressures.⁴⁹

For the double-decoupling simulations, solvent moves were attempted with a probability of 85.7%, protein side-chain moves with a probability of 12.9%, and solute moves with a probability of 1.4%. As with the JAWS simulations, only the bond angles and torsions for the side chains of residues within 10 Å of any heavy atom of zanamivir were sampled.

Error estimates from the double-decoupling simulations were obtained as the standard error across at least three independent simulations.

GCMC Protocol: Interacting Particle Method. The GCMC simulations for the individual water molecules were performed using the interacting particle method.⁶¹ Insertion and deletion attempts of water molecules were accepted using the following Metropolis tests.

$$P_{\rm in} = \min \left[1, \, \frac{\exp(B)}{N+1} \exp\left(\frac{-\Delta E}{k_{\rm B} T}\right) \right] \tag{7}$$

$$P_{\text{del}} = \min \left[1, N \exp(-B) \exp\left(\frac{-\Delta E}{k_{\text{B}} T}\right) \right]$$
 (8)

In the above equations, N is the number of particles in the simulation and B is the Adams parameter ($B = \mu'/k_{\rm B}T + \ln < N >$). Here < N > is the average number of particles in the system at the specific B value at which the simulation is run. ²⁹ μ' is the excess chemical potential, and ΔE is the change in energy between the new and old states.

In the GCMC simulations, each water molecule was looked at individually in this study to calculate its binding free energy; calculating the binding free energy of all of the waters within the same simulation is a demanding task which will result in sampling difficulties over short simulation lengths. Consequently, each water was allocated its own defined GCMC simulation region, where other water molecules were prohibited from entering the GCMC region using a hardwall but solute and protein atoms were allowed to occupy the same region. A smaller 2 Å \times 2 Å \times 2 Å grid was defined around each water molecule to obtain sufficient sampling of the localized water occupancy, since it was observed that in some cases a larger volume occupied by the water molecule was filled with a solute atom, meaning that poor acceptance rates were observed.

Each B-value was simulated for 40 million MC moves, divided into 800 blocks of 50 000 steps each. At the end of each simulation the average water population across the entire simulation was recorded. The decoupling free energy of the water was found using eq 9.

$$\Delta G_{\text{dec}} = -k_{\text{B}}T \ln \left(\frac{[L_{\text{sim}}]}{[L_{\text{ideal}}]} \right)$$
(9)

In eq 9, $[L_{\text{sim}}]$ is found by recording the population at a particular B value. This population is converted into a localized concentration by dividing by the simulation volume and, then, converting this into a molar concentration using Avogadro's number. $[L_{\text{ideal}}]$, the concentration of an ideal gas in the simulation

volume, is related to the *B* value of the simulation and is found using eq 10.

$$[L_{\text{ideal}}] = 55.56 \text{Mexp B} \tag{10}$$

Having calculated $\Delta G_{\text{de}\sigma}$ the binding free energy of the water was found using eq 11.

$$\Delta G_{\rm bind} = \Delta G_{\rm dec} - \Delta G_{\rm hyd} \tag{11}$$

For each water molecule, at least four *B* values were simulated to allow for a reliable estimate of the binding free energy, found as the average of the binding free energies across the range of *B* values. *B* values which gave <*N*> greater than 0.8 and less than 0.1 were not used in the analysis.

For the GCMC simulations, solvent moves were attempted with a probability of 44%, protein side-chain moves with a probability of 5.8%, and solute moves with a probability of 1.8%. Insertion and deletions were each attempted with an equal probability of 2.2%, with translations and rotations of the isolated GCMC water attempted with a probability of 44%.

Using the interacting particle approach, an estimate of the error in the binding free energy of a water can be found as standard error of the binding free energy across a number of independent simulations performed with different *B* values.

GCMC Protocol: Simulated Annealing Method. An alternative method for calculating the binding free energy of a water molecule through GCMC lies in the simulated annealing approach. Rather than converting populations into localized concentrations, the populations obtained from simulating at a range of B values are instead used to make a free energy titration plot. The value of B can be related to the water binding free energy using eq 12.

$$\Delta G_{\text{bind}} = k_{\text{B}} T(B - \ln \langle N \rangle) - \Delta G_{\text{hyd}}$$
(12)

By plotting the average population of the water molecule as a function of the binding free energy, found from B, the value of $\Delta G_{\rm bind}$ can be estimated as the binding free energy observed when the site is mostly empty.

Using the simulated annealing approach, an estimate of the error in the binding free energy of a water can be found as the standard error of the population across a number of independent simulations performed at the same *B* value and observing the subsequent range in the binding free energy.

The simulation protocol for the simulated annealing approach is the same as for the interacting particle method.

Unless stated otherwise, the above protocols were used for all of the subsequent studies. The JAWS and GCMC protocols were coded into the in-house Monte Carlo software, ProtoMS, 59 which was used to perform all the simulations.

RESULTS AND DISCUSSION

JAWS Placement. A JAWS stage one simulation was performed upon N9-neuraminidase, incorporating a grid volume of 1053 Å³. The native crystal structure contains a network of five crystallographic waters with two of these, Wat567 (subsequently referred to as Wat5) and Wat568 (subsequently referred to as Wat4), forming direct interactions with each other, the protein, and the zanamivir ligand. Wat4 is stabilized by a triad of nearby waters; Wat508 (Wat3), Wat543 (Wat2) and Wat575 (Wat1), which are all hydrogen-bonded to both each other and the protein. An additional structural water

molecule, Wat507 (Wat6), is found in an isolated cavity and has no direct interactions with the zanamivir ligand. The JAWS simulation identified seven possible hydration sites, shown in Figure 1, in good agreement with both the crystallographic data and the original simulations performed by Michel et al.⁴¹

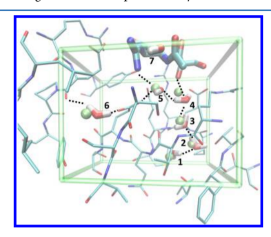


Figure 1. Seven possible hydration sites identified by JAWS stage 1 in N9 neuraminidase. Crystallographic waters are shown in light green, with critical hydrogen bonds involving the protein, zanamivir, and JAWS-waters shown as dashed lines. The zanamivir ligand is shown in licorice.

Attempts were made to calculate the binding affinity of each of the waters using the JAWS stage 2 algorithm. It was found, however, that the majority of water molecules did not experience sufficient θ < 0.05 transitions during the simulation time frame. As a result, the binding free energies calculated by eq 3 were either poorly converged or unavailable. An example of this is shown in Figure 2, where the standard biasing term

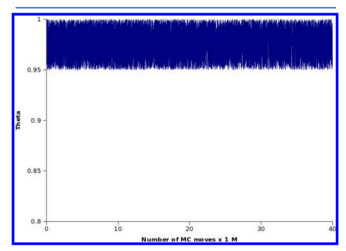


Figure 2. θ Sampling for Wat5 in N9 neuraminidase using a biasing potential of 6.4 kcal mol⁻¹.

does not induce any θ < 0.05 transitions for Wat 5 in N9 neuraminidase: the water is always on in the simulation, and hence, the free energy cannot be estimated using eq 3.

It has been previously recognized that one the major drawbacks of the JAWS algorithm is that it cannot calculate the binding affinities of strongly bound waters. To calculate the binding free energies of strongly bound water molecules, we have modified the JAWS biasing term. These modifications are now discussed.

Extension of the JAWS Algorithm to Calculate $\Delta G_{\rm bind}$ for Strongly Bound Waters. The calculation of the binding free energy of a water molecule is captured by eq 3. For weakly bound water molecules, the biasing potential applied in the second stage of the JAWS algorithm is sufficient to ensure that the θ water molecule can sample both the on and off states, ensuring that enough statistical sampling is performed to obtain a reliable free energy estimate. However, for strongly bound water molecules, the bias potential used in previous studies is not sufficient to induce transitions to the off state, resulting in either poor or no sampling and an unreliable estimate of the binding free energy.

One way of ensuring that sufficient sampling is performed at both end states is by changing the biasing potential applied in the second stage of the algorithm. Rather than basing this upon the hydration free energy of water, the applied bias can be changed to one which induces sufficient transitions between the two states. This is achieved by replacing $-\Delta G_{\rm hyd}$ in eq 1 with $\Delta G_{\rm bias}$. The resulting free energies obtained are indicative of the new biasing potential and, hence, must be corrected to take this into account, as shown in eq 13 where $\Delta G_{\rm bias}$ is the value of the bias applied in the second stage of the algorithm.

$$\Delta G_{\text{bind}}(\text{water, site } i) = -k_{\text{B}}T \ln \left(\frac{P(\theta_i \to 1)}{P(\theta_i \to 0)}\right) - \Delta G_{\text{hyd}} - \Delta G_{\text{bias}}$$
(13)

The use of a threshold to define the on and off states induces an error in the calculated values using eq 13, since the thresholds for the end states are taken to be 0.95 and 0.05, respectively, rather than 1 and 0. Since the chance of observing a water molecule being completely turned on $(\theta=1)$ or completely off $(\theta=0)$ is unlikely during the simulation time frame, an approximation is made to define the on and off states. Assuming that the water population distribution of θ across 0.95–1.00 and 0.05–0.00 is uniform, eq 13 can be modified to eqs 14 and 15. In these equations, the biasing correction is approximated by the average of the applied threshold and the absolute end-points $(\theta=1)$ and $(\theta=0)$:

$$\Delta G_{\text{corr}} = 0.975 \Delta G_{\text{bias}} - 0.025 \Delta G_{\text{bias}}$$
 (14)

$$\Delta G_{\rm bind}(\text{water, site } i) = -k_{\rm B}T \ln \left(\frac{P(\theta_i \to 1)}{P(\theta_i \to 0)}\right) - \Delta G_{\rm hyd} - \Delta G_{\rm corr}$$
(15)

The free energies obtained by this method are broadly independent of the applied bias, providing θ > 0.95 and θ < 0.05 are adequately sampled, as shown in Figure 3.

The free energies shown in Figure 3 were found using eq 15, with the associated error found as

$$\Delta G_{\text{error}} = \Delta G_{\text{bias}} - \Delta G_{\text{corr}}$$
 (16)

The need for changing the applied bias can be seen in Figure 4, whereby transitions between the on and off state are induced by increasing the applied bias. At a biasing potential, $\Delta G_{\rm bias}$, of 10 kcal ${\rm mol}^{-1}$, the water molecule experiences most of the simulation time in the on state, with no transitions to the off state being observed, meaning that a reliable free energy estimate cannot be obtained. However, upon using a $\Delta G_{\rm bias}$ of 17 kcal ${\rm mol}^{-1}$, the water molecule can sample both end states, allowing a reliable free energy estimate to be obtained.

There is also an error associated with the choice of θ threshold. For example, the threshold for an "on" state could arbitrarily be either $\theta > 0.95$ or $\theta > 0.98$. To estimate the associated error,

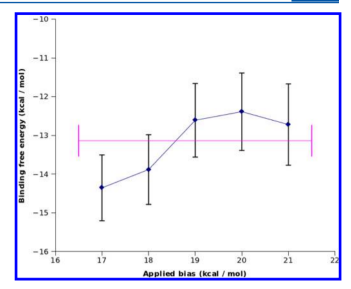


Figure 3. Effect of the applied biasing potential upon the threshold-corrected JAWS stage 2 binding free energy of Wat5 in N9 neuraminidase. The average binding free energy across the five simulations is shown in magenta.

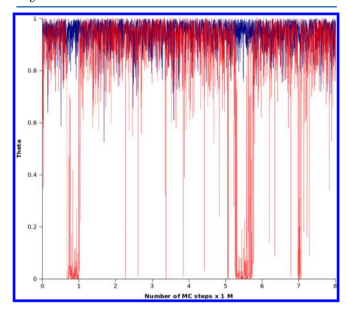


Figure 4. θ Sampling as a function of the applied bias potential for Wat3 in N9 neuraminidase. Shown in blue is the sampling at a bias of 10 kcal mol⁻¹, while the sampling at 17 kcal mol⁻¹ is shown in red.

the binding free energy of Wat5 in N9-neuraminidase was calculated using different thresholds for both the on and off states. The calculated free energies can be found in Figure 5, with the error associated with the choice of θ , found as the standard error across the five measurements, estimated to be ± 0.40 kcal mol⁻¹. It can be seen that there seems to be no systematic dependence on the theta threshold chosen.

The overall error for the JAWS simulations is thus the combination of the error in the θ threshold and eq 16.

Choice of Biasing Potential. Since the binding free energy is broadly independent of the applied bias, provided the on and off states are adequately sampled, a JAWS stage 2 simulation needs to be run at only one value of the bias to extract the binding free energy. The ideal bias should induce an equal number of on and off states, meaning that the binding free energy becomes the difference between the standard hydration

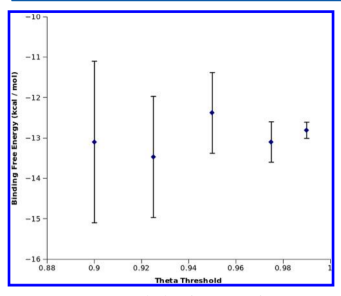


Figure 5. JAWS stage 2 binding free energy of Wat5 in N9 neuraminidase as a function of the θ threshold applied. Simulations were performed using a biasing potential of 20 kcal mol⁻¹, with errors found using eq 16.

free energy of water and the applied bias, as in eq 15. An equal number of on and off states means that the water is, on average, present 50% of the time and should give the most reliable estimate of the binding free energy. To achieve this, a simple optimization procedure of the biasing potential has been applied.

A short JAWS stage 2 simulation, typically one million MC moves, is performed and an estimation of the binding free energy found. The biasing potential is then optimized to obtain a value of the bias which yields an equal number of on and off states. As an example, the process is illustrated for hydration site Wat3 in Table 1.

Table 1. Optimization Procedure for Wat3 in N9 Neuraminidase

iteration	$\Delta G_{ ext{bias}}$ (kcal mol ⁻¹)	ln(on/ off)	iteration	$\Delta G_{ m bias} \ ({ m kcal \ mol^{-1}})$	ln(on/ off)
1	15.0	12	6	17.5	-1.0
2	15.5	10	7	17.0	11
3	16.0	11	8	17.5	3.0
4	16.5	4.0	9	18.0	-2.0
5	17.0	3.0	10	17.5	0.0

In Table 1, the iterative optimization is trying to obtain a value of ΔG_{bias} which induces an equal number of on and states. As shown in eq 15, an equal number of on and off states should result in the log term approaching zero. At the end of each simulation, the log ratio of on and off states is calculated. If the value is positive, suggesting more on states than off, then the value of ΔG_{bias} is increased by 0.5 kcal mol^{-1} for the next iteration. If the value is negative, suggesting more off states than on, then the value of ΔG_{bias} is decreased by 0.5 kcal mol⁻¹. At the end of the process, typically incorporating 10 iterations, the value of ΔG_{bias} which gives a log term of zero is chosen for the main simulation. In this example, the value of ΔG_{bias} was taken to be 17.5 kcal mol⁻¹. The variability in the calculated log terms shows that this is an approximate method and that longer runs are required to achieve convergence, but they provide guidance to the most suitable value of ΔG_{bias} .

■ BINDING FREE ENERGY CALCULATIONS

Comparison of JAWS and RETI. Using the new modifications, a JAWS stage 2 simulation was performed upon each hydration site as identified in Figure 1, with each site also studied using double-decoupling. The binding free energy for JAWS stage two simulations was found using eq 15. The free energy comparison between the two methods can be seen in Figure 6.

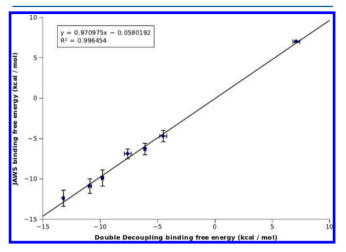


Figure 6. Binding free energies for the seven hydration sites in N9 neuraminidase, found using JAWS stage 2 and RETI double-decoupling.

Figure 6 clearly demonstrates that the two methods give excellent agreement with each other. Binding affinities of both strongly and weakly bound water molecules are calculated by the two methods, showing that the modification to the JAWS algorithm has been successful in predicting the binding free energy of water molecules which previously were incalculable. The biasing potentials used to calculate the JAWS free energies in Figure 6 are given in Table 2.

Table 2. Binding Free Energies for the Seven Water Molecules in N9 Neuraminidase, Found Using JAWS Stage 2 and RETI Double-Decoupling a

water molecule	JAWS bias (kcal mol ⁻¹)	JAWS ΔG_{bind} (kcal mol ⁻¹)	RETI $\Delta G_{ ext{bind}}$ (kcal mol ⁻¹)
1	14	-4.7(0.7)	-4.5 (0.3)
2	12	-6.9 (0.6)	-7.6(0.3)
3	17.5	-10.9(0.9)	-10.9(0.1)
4	14	-6.3(0.7)	-6.1(0.1)
5	20	-12.4 (1.0)	-13.2(0.1)
6	19	-9.9 (1.0)	-9.8 (0.1)
7	2	7.0 (0.1)	7.1 (0.3)

^aStatistical errors are shown in parentheses. The individual free energy components for these double-decoupling simulations can be found in the Supporting Information.

The convergence for the JAWS stage 2 simulations as a function of the simulation length are given in Figure 7. It can be seen that all of the simulations are well converged after ca. 10 million MC moves, demonstrating further that the biasing potentials used yield precise results.

Comparison of GCMC and RETI. Using the hydration sites identified by the JAWS stage 1 simulations, the free energy of binding of each site was calculated using the interacting particle

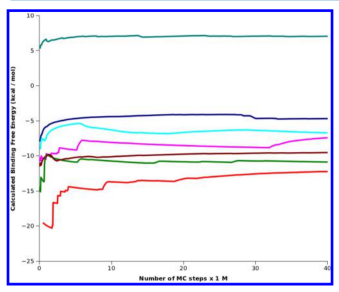


Figure 7. Convergence of the binding free energies in the JAWS stage 2 simulations. Key: Wat1 (dark blue), Wat2 (magenta), Wat3 (dark green), Wat4 (cyan), Wat5 (red), Wat6 (maroon), Wat7 (turquoise).

method of Clark et al.⁶¹ Equation 9 yields the decoupling energy of the water molecule from the protein and can be corrected with the hydration free energy of water to arrive at a binding free energy using eq 11. Figure 8 shows the predicted

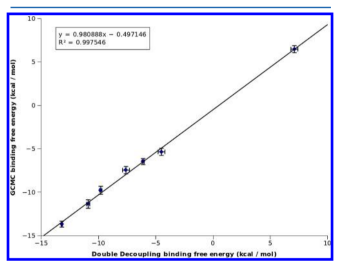


Figure 8. Binding free energies for the seven hydration sites in N9 neuraminidase, found using GCMC and RETI double-decoupling.

binding affinity of the seven molecules using the two methods and shows an excellent correlation.

The reported GCMC binding free energies were calculated as the average of the binding free energy across a range of B values. An example of this for Wat7, the weakest binder in the series, can be seen in Table 3. The table shows that the calculated binding free energy is consistent and approximately 6.5 kcal mol^{-1} once the average population recorded throughout the simulation, used to calculate $[L_{\mathrm{sim}}]$, drops below 0.8 and is above 0.1, corresponding to sufficient sampling of the water populations. This behavior is consistent with that demonstrated by Clark et al. in the calculation of benzene—T4 lysozyme binding free energies. 61,62 The maximum occupancy for the water site is unity, meaning statistically significant occupancies of less than one need to

Table 3. Calculated Free Energies for Wat7 in N9 Neuraminidase, Found at Different B Levels

В	average number of molecules	$egin{aligned} [L_{ ext{sim}}] \ (M) \end{aligned}$	$egin{pmatrix} [L_{ m ideal}] \ (M) \end{pmatrix}$	$\Delta G_{ m dec}$ (kcal mol ⁻¹)	$\Delta G_{\rm bind} \ ({ m kcal \ mol^{-1}})$
4	0.97	200	3030	1.6	8.0
3	0.93	190	1120	1.0	7.4
2	0.73	150	410	0.6	7.0
1	0.57	120	150	0.1	6.5
0	0.37	75	56	-0.2	6.2
-1	0.15	30	20	-0.2	6.2
-2	0.04	8	8	0.0	6.4

be obtained to calculate a reliable estimate of the binding free energy using eq 9.

Using the data for *B* values less than 3 and greater than -2 in Table 3, the binding free energy can be estimated as $6.48 \pm 0.19 \text{ kcal mol}^{-1}$.

One major potential drawback associated with the GCMC method lies in the acceptance rate of insertion and deletion moves. For an insertion to be accepted it is important that the orientation of the water molecule is correct, since otherwise it is likely that the intermolecular interactions between the water and its environment will be unfavorable.³⁴ As a result, insertion rates as low as 0.1% are seen in GCMC simulations, which in turn leads to poor sampling. It is this poor sampling which could potentially lead to an increase in the uncertainty in the free energies derived from GCMC simulations compared to both double-decoupling and JAWS, but no evidence of this has been seen in this study. Although an insertion rate of 0.1% at first sight is worryingly low, this corresponds to successfully sampling 800 insertion events over the simulation, which appear sufficient to converge the free energy estimate.

Since the interacting particle method can generate populations as a function of B, this information can also be used to derive free energies via the simulated annealing approach. The data from Table 3 have been used to generate such a titration profile, seen in Figure 9. Values of $\langle N \rangle$ below 0.1 were not used in the analysis since, for low values of $\langle N \rangle$, small variations can have a large effect on the logarithmic term and make the binding free energy estimate imprecise.

Figure 9 shows that the estimated binding free energy of Wat7 is 6.9 ± 0.3 kcal mol⁻¹, in reasonable agreement with the value calculated by the interacting particle method. Since either method can be used to derive broadly the same result to within error, the question arises as to which of the GCMC methods is to be preferred when calculating binding free energies. While the simulated annealing approach gives information regarding the behavior of the system as a function of B, the interacting particle approach is significantly faster since it only requires the simulation to be performed at one value of B. However, the optimal B value to choose is not always known a priori, meaning that it can require several different simulations to identify a B value that yields precise binding free energies. It therefore appears to be advantageous to use the simulated annealing approach to see how the population changes as a function of the applied B value, and simultaneously to use the interacting particle method to calculate the binding free energy.

Method Comparison. The N9-neuraminidase system was chosen as a test case for the different free energy methodologies since there are a large number of studies which have used the system in free energy calculations. ^{41,49} The fact that all three free-energy methods give near identical results, to within

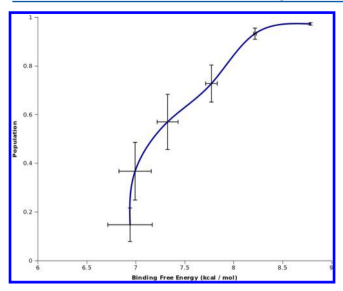


Figure 9. Free energy titration plot for Wat7 in N9 neuraminidase, found using the GCMC simulated annealing approach. The population error bars were found as the standard error across two independent simulations, with this error used to calculate the associated error in the binding free energy using eq 12.

statistical error, is clearly encouraging and lends itself to the question of which method is best suited to a particular problem. While RETI double-decoupling is the most rigorous method of the three, it is also the most computationally expensive and requires knowledge of the water binding sites. A typical simulation requires in excess of 300 CPU h on 16 2.6 GHz processors, while both GCMC and JAWS require an order of magnitude less computational time and can be run on a single processor. As a result it is suggested that double decoupling is used in cases where precise free energies are required.

One drawback of the double-decoupling approach is that it requires prior knowledge of the water binding positions; something which is found dynamically in both JAWS and GCMC. As such, if systems without clearly defined crystallographic waters are studied then either JAWS, GCMC, or both methods should be employed to identify potential hydration sites. JAWS has already been employed in free energy studies, whereby changes in hydration as a function of ligand perturbations were evaluated. With the extension of the JAWS biasing potential, both JAWS and GCMC can calculate the binding free energy for both strongly and weakly bound waters during a single simulation and take similar simulation times. However, one possible advantage in the JAWS approach is that once the optimal biasing potential is found no further simulations need to be run while the GCMC approach requires several simulations at different B values to arrive at the binding free energy. This in itself, however, highlights one of the advantages in the GCMC approach; that it can give information on the binding of molecules as a function of the chemical potential.

One potential problem with the JAWS approach to calculate binding free energies is how to deal with the intermediate θ states. Since only the end points are used when calculating the binding free energy, it is unclear whether or not the intermediate data should be considered. The probability that a θ -water adopts an ill-defined intermediate state can be significant, as indicated in Figure 10.

Upon the basis of the excellent agreement between JAWS, GCMC, and double-decoupling, it seems that ignoring the

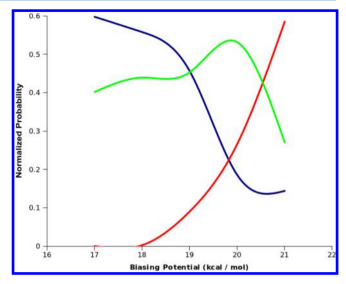


Figure 10. Normalized sampling probabilities for the on, off, and intermediate states as a function of the applied biasing potential for Wat 5 in N9 neuraminidase. The on state (dark blue) was defined as $\theta > 0.95$, the off state (red) was defined as $\theta < 0.05$, and the intermediate state (green) accounted for the remaining θ values.

intermediate states is acceptable. This is, however, a waste of potentially useful data, although it is unclear how the data could be analyzed.

CONCLUSIONS

The application of JAWS, GCMC, and double-decoupling Monte Carlo has demonstrated that all three methods yield water binding free energies which are consistent with each other when applied to a single, isolated water site in N9 neuraminidase. The double-decoupling approach is by far the slowest of the three methods, since it requires free energy changes for intermediate states between $\lambda = 0$ and 1 to be simulated. The method is the most rigorous of the three studied, and we propose that it is used to score water molecules when precise free energies are required.

The extended JAWS stage 2 and GCMC simulations both require, on average, an order of magnitude less computational resources than the double-decoupling method, meaning they should be preferred for rapid estimation of the binding free energy of a single water molecule. Despite this, the JAWS methodology typically requires fewer simulations to be run than GCMC, and on this basis we propose that JAWS should be preferred in the calculation of water binding free energies.

Although both JAWS and GCMC are capable of predicting the location of water molecules in protein cavities, the resulting data from a JAWS simulation can be difficult to interpret. For welldefined sites, such as the N9-neuraminidase system, presented in this paper, the population maps are clear and easy to interpret. However, if the population density contours overlap, it can prove difficult to resolve the population densities into individual hydration sites. The population density does not reveal whether the binding of two or more water molecules are correlated, or whether a single water molecule hops between two adjacent sites. In comparison the GCMC method is capable of clearly identifying hydration sites, since a representative simulation snapshot with the desired number of molecules can be easily extracted. Crucially, none of the GCMC snapshots contain illdefined intermediate θ -water states and each snapshot is a valid structural representation for the particular binding free energy at which the simulation was run. As such, cooperative binding between water molecules can also be identified, something which is harder to infer from a JAWS population density analysis.

In cases where the JAWS density map does not indicate clear, isolated water binding sites, we propose that GCMC should be used to identify potential hydration sites. Since the results from a GCMC simulation are dependent upon the chemical potential used, the chemical potential can be tuned to observe hydration patterns as a function of the binding free energy. The resulting hydration sites could then be scored in a JAWS stage 2 simulation if desired.

ASSOCIATED CONTENT

S Supporting Information

Protonation state analysis for the 1nnc protein structure and partial charges used to model the zanamivir ligand. Free energy decomposition for the decoupling of water molecules bound to 1nnc and zanamivir using RETI double-decoupling. This material is available free of charge via the Internet at http://pubs.acs.org/.

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Notes

The authors declare no competing financial interest.

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