Computing Clinically Relevant Binding Free Energies of HIV-1 Protease Inhibitors

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ABSTRACT: The use of molecular simulation to estimate the strength of macromolecular binding free energies is becoming increasingly widespread, with goals ranging from lead optimization and enrichment in drug discovery to personalizing or stratifying treatment regimes. In order to realize the potential of such approaches to predict new results, not merely to explain previous experimental findings, it is necessary that the methods used are reliable and accurate, and that their limitations are thoroughly understood. However, the computational cost of atomistic simulation techniques such as molecular dynamics (MD) has meant that until recently little work has focused on validating and verifying the available free energy methodologies, with the consequence that many of the results published in the literature are not reproducible. Here, we present a detailed analysis of two of the most popular approximate methods for calculating binding free energies from molecular simulations, molecular mechanics Poisson–Boltzmann surface area (MMPBSA) and molecular mechanics generalized Born surface area (MMGBSA), applied to the nine FDA-approved HIV-1 protease inhibitors. Our results show that the values obtained from replica simulations of the same protease–drug complex, differing only in initially assigned atom velocities, can vary by as much as 10 kcal mol⁻¹, which is greater than the difference between the best and worst binding inhibitors under investigation. Despite this, analysis of ensembles of simulations producing 50 trajectories of 4 ns duration leads to well converged free energy estimates. For seven inhibitors, we find that with correctly converged normal mode estimates of the configurational entropy, we can correctly distinguish inhibitors in agreement with experimental data for both the MMPBSA and MMGBSA methods and thus have the ability to rank the efficacy of binding of this selection of drugs to the protease (no account is made for free energy penalties associated with protein distortion leading to the over estimation of the binding strength of the two largest inhibitors ritonavir and atazanavir). We obtain improved rankings and estimates of the relative binding strengths of the drugs by using a novel combination of MMPBSA/MMGBSA with normal mode entropy estimates and the free energy of association calculated directly from simulation trajectories. Our work provides a thorough assessment of what is required to produce converged and hence reliable free energies for protein–ligand binding.

1. INTRODUCTION

In the past two decades, both computationally assisted rational drug design and personalized medicine have been held out as potential new paradigms which would produce novel therapies at drug design and personalized medicine have been held out as potential new paradigms which would produce novel therapies. In order to realize the potential of such approaches to predict new results, not merely to explain previous experimental findings, it is necessary that the methods used are reliable and accurate, and that their limitations are thoroughly understood. However, the computational cost of atomistic simulation techniques such as molecular dynamics (MD) has meant that until recently little work has focused on validating and verifying the available free energy methodologies, with the consequence that many of the results published in the literature are not reproducible. Here, we present a detailed analysis of two of the most popular approximate methods for calculating binding free energies from molecular simulations, molecular mechanics Poisson–Boltzmann surface area (MMPBSA) and molecular mechanics generalized Born surface area (MMGBSA), applied to the nine FDA-approved HIV-1 protease inhibitors. Our results show that the values obtained from replica simulations of the same protease–drug complex, differing only in initially assigned atom velocities, can vary by as much as 10 kcal mol⁻¹, which is greater than the difference between the best and worst binding inhibitors under investigation. Despite this, analysis of ensembles of simulations producing 50 trajectories of 4 ns duration leads to well converged free energy estimates. For seven inhibitors, we find that with correctly converged normal mode estimates of the configurational entropy, we can correctly distinguish inhibitors in agreement with experimental data for both the MMPBSA and MMGBSA methods and thus have the ability to rank the efficacy of binding of this selection of drugs to the protease (no account is made for free energy penalties associated with protein distortion leading to the over estimation of the binding strength of the two largest inhibitors ritonavir and atazanavir). We obtain improved rankings and estimates of the relative binding strengths of the drugs by using a novel combination of MMPBSA/MMGBSA with normal mode entropy estimates and the free energy of association calculated directly from simulation trajectories. Our work provides a thorough assessment of what is required to produce converged and hence reliable free energies for protein–ligand binding.

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large numbers of candidates must be assessed, and clinical applications, where diagnoses must typically be given in days or at most weeks, rapid turnaround of calculations is vital, and calculations that take a week or more to complete are unlikely to ever be adopted beyond basic research environments. More fundamentally, it is not the notional accuracy of the different methods but the reproducibility of the results obtained which needs to be verified and validated before any method can be seen as a reliable tool for future applications.

The molecular mechanics Poisson–Boltzmann surface area (MMPBSA)\(^{10,11}\) and molecular mechanics generalized Born surface area (MMGBSA)\(^{12}\) methods of estimating binding free energies have become some of the most widely applied due to their claims to provide a compromise between accuracy and speed. Both of these methods, however, neglect to account for changes in configurational entropy upon binding. Consequently, they are frequently combined with estimates of this contribution made using normal-mode analysis of harmonic frequencies from minimized snapshots from the MD trajectory. While the convergence and applicability of MMPBSA and MMGBSA have been widely studied\(^{13,14}\), the computational cost of normal-mode analysis has meant that only recently has its performance begun to be similarly investigated.\(^{15}\) Here, we present an extensive analysis of the convergence properties and ability to reproduce experimental values of the methods applied to HIV-1 protease binding to the nine FDA-approved inhibitors. These results build on our previous work that used these techniques to evaluate differences in binding strength of different protein sequences to individual drugs in both the HIV protease\(^{16,17}\) and the anticancer therapy target epidermal growth factor receptor (EGFR).\(^{18}\) In the latter case, the prediction made for the L858R mutant was thought to disagree with experimental results,\(^{19}\) but more recent data have confirmed the validity of the computational estimates.\(^{20}\)

Ensembles of multiple short simulations were run for each protein–ligand combination, as previous studies have suggested that a multiple/ensemble MD approach can sample phase space more efficiently than fewer long time simulations.\(^{16,21,22}\) This approach also allows us to investigate the reproducibility of the calculations on the individual simulations within an ensemble (we will refer to an individual simulation within an ensemble for each inhibitor as a replica). We also evaluate the performance of a more recent data have confirmed the validity of the computational estimates.

2. METHODS

When two reactants combine at constant temperature and pressure, the binding affinity is characterized by the change in Gibbs free energy given by

\[
\Delta G = \Delta H - T \Delta S
\]  

(1)

at temperature \(T\) and is composed of the change in enthalpy (\(\Delta H\)) and the change in entropy (\(\Delta S\)). In this study, we aim to assess the ability of several different end-point free energy calculation methodologies to estimate \(\Delta G\) using protein–ligand conformations generated via molecular dynamics simulations. The methods under investigation are MMPBSA\(^{10,11}\) and MMGBSA,\(^{12}\) including and excluding terms aiming to explicitly incorporate solute entropic contributions to the binding energy. All of these methods run as post-processing steps on snapshots taken from the MD trajectories, with averages computed over all snapshots.

Even when incorporating terms describing solute entropic contributions, these methods possess several apparent limitations for computing absolute binding free energies. They do not implicitly account for free energy differences that arise from possible changes in key protonation states or explicit water-mediated binding between a protein and ligand which may provide significant contributions to the binding free energy. Despite these limitations and scepticism about the validity of the methods in some quarters,\(^{23}\) our previous work indicates that changes in binding energy of less than 1 kcal mol\(^{-1}\) between HIV-1 protease mutants can be distinguished.\(^{16,26}\)

2.1. Free Energy Calculation Protocols. Both MMPBSA and MMGBSA are end-point free-energy calculations. In such methods, the free energy of binding, \(\Delta G\), is calculated using

\[
\Delta G = \langle G_{\text{complex}} \rangle - \langle G_{\text{enzyme}} \rangle - \langle G_{\text{ligand}} \rangle
\]

(2)

where \(\langle G_{\text{complex}} \rangle\), \(\langle G_{\text{enzyme}} \rangle\), and \(\langle G_{\text{ligand}} \rangle\) are the average values of the Gibbs free energy for the complex, enzyme, and ligand, respectively. Separate simulation of the complex and its two components can be used, but due to difficulties in obtaining converged free energies, it is more common to extract configurations of the free enzyme and ligand from simulation of the complex. Much of the improved convergence behavior of the single trajectory approach is due to the fact that the noisy term describing the internal energy of the system components is canceled between ligand, enzyme, and complex.\(^{13}\) This is the strategy we have employed in the present study. Normal mode solute entropy calculations use the same configurations and also a calculation similar to eq 2 to calculate the overall entropic penalty from values computed for the complex, enzyme, and ligand. The free energy of association, however, deals exclusively with the changes in binding energy of less than 1 kcal mol\(^{-1}\) and is computed from collections of snapshots. The details of all four computations are provided below.

2.1.1. MMPBSA and MMGBSA. The binding free energy change calculated by MMPBSA and MMGBSA (\(\Delta G_{\text{MMPBSA/GBSASSA}}\)) can be broken down into a number of components:

\[
\Delta G_{\text{MMPBSA/GBSASSA}} = \Delta G_{\text{MM}}^{\text{ele}} + \Delta G_{\text{vdW}}^{\text{MM}} + \Delta G_{\text{int}}^{\text{MM}} + \Delta G_{\text{nonpol}}^{\text{MM}} + \Delta G_{\text{pol}}^{\text{sol}}
\]

(3)

where \(\Delta G_{\text{MM}}^{\text{ele}}\) and \(\Delta G_{\text{MM}}^{\text{vdW}}\) are the van der Waals and electrostatic contributions to the molecular mechanics free energy difference, respectively, and \(\Delta G_{\text{pol}}^{\text{sol}}\) and \(\Delta G_{\text{nonpol}}^{\text{pol}}\) are the polar and nonpolar solvation terms, respectively. Modules of the AMBER 9 package\(^{27}\) were used in the evaluation of all components of the MMPBSA calculation. The SANDER module was employed to calculate both molecular mechanics terms (\(\Delta G_{\text{vdW}}^{\text{MM}}\) and \(\Delta G_{\text{ele}}^{\text{MM}}\)), with no cutoff being applied to the nonbonded energies. The electrostatic free energy of solvation, \(\Delta G_{\text{pol}}^{\text{sol}}\) is the part of the calculation described by the Poisson–Boltzmann (PB) or generalized Born (GB) calculation. GB is an approximation to PB, and in both methods internal and external dielectric constants of 1 and 80, respectively, were used. In the PB case, a thousand iterations of the linear Poisson–Boltzmann equation were performed on a cubic lattice grid with a spacing of 0.5 Å using the DelPhi\(^{28}\) program. The GB calculations were calculated using the model proposed by Onufriev et al.\(^{12}\) The nonpolar
solvent energy, \( \Delta G_{\text{nonpol}}^{\text{sol}} \) was calculated from the solvent accessible surface area (SASA) using the MSMS program\(^{29} \) with a 1.4 Å radius probe based on the equation:

\[
\Delta G_{\text{nonpol}} = \gamma A + \beta
\]

where \( A \) is the solvent accessible surface area difference, \( \gamma \) is the surface tension, and \( \beta \) is the offset. The constants \( \gamma \) and \( \beta \) were set to the standard values of 0.0052 kcal mol\(^{-1}\) Å\(^{-2}\) and 0.92 kcal mol\(^{-1}\), respectively, in our MMPBSA simulations and 0.0072 kcal mol\(^{-1}\) Å\(^{-2}\) and 0 kcal mol\(^{-1}\) in MMGBSA\(^{10-12} \).

2.1.2. Normal Mode Analysis. The non-polar solvation term of the MM(GB)PB calculation incorporates an implicit estimate of the entropic changes associated with the insertion of a solute into the solvent. However, no account is made for the entropic contribution of the entropic changes associated with the insertion of a solute into the solvent. However, no account is made for the entropic penalty of the binding reaction.

\[
\Delta G_{\text{Assoc}} = -RT \ln \left( \frac{C^0_{\text{trans rot}} z_{\text{lig}}^{\text{rot}}}{8\pi^2} \right)
\]

where \( C^0 \) is the standard state concentration, usually taken to be 1 M (1 molecule/1660 Å\(^3\)), \( z_{\text{lig}}^{\text{rot}} \) and \( z_{\text{lig}}^{\text{rot}} \) are the ligand translational and rotational configuration integrals, respectively. \( z_{\text{lig}}^{\text{trans}} \) was calculated by superimposing every snapshot onto an average structure of the protease using Cα atoms alone. This provides a static reference system along with an average ligand structure. The eigenvalues \( \lambda_i \) of the covariance matrix of the ligand center of mass were then computed providing variance measures along three independent axes from the relation \( \lambda_i = \langle \Delta x_i^2 \rangle \). The translational configurational integral is then given by

\[
z_{\text{lig}}^{\text{rot}} = (2\pi)^{3/2} \left( \langle \Delta x_1^2 \rangle \langle \Delta x_2^2 \rangle \langle \Delta x_3^2 \rangle \right)^{1/2}
\]

Similarly, the ligand’s rotational integral can be computed from quaternions representing the ligand’s rotational motion relative to the average structure. A small angle approximation allows the reduction of the three quaternions describing rotation about three axes into a single quaternion, three of whose components are sinusoidally related to three rotation angles. As for the translational component, the covariance matrix can be evaluated to produce eigenvalues that can be used to calculate \( z_{\text{lig}}^{\text{rot}} \) in an analogous manner to eq 10.

The structural superposition of each snapshot onto the average structure was performed using scripts based on the alignment code found in MDAnalysis.\(^{31-33} \) The use of this method is attractive as it is based on the configurations taken directly from the simulations and is computationally inexpensive. This method assumes that the simulations explore a single minimum, whereas we expect that replicas will explore different minima. Consequently, here we evaluate \( \Delta G_{\text{Assoc}} \) for each replica individually.

2.2. Model Preparation and Simulation Protocol. Preparation and simulation setup were performed using the automated Binding Affinity Calculator (BAC); full details of this tool and the simulation parameters employed are available in Sadiq et al.\(^ {34} \) Models of the subtype B HXB2 wildtype sequence of HIV-1 were constructed using the coordinates from PDB crystal structures listed in Table 1. All systems were solvated in orthorhombic water boxes with a minimum extension from the protein of 14 Å.

Protein parameters were taken from the standard AMBER force field for bioorganic systems (ff03).\(^ {35} \) Drug parameters were produced using the general AMBER force field (GAFF)\(^ {36} \) following the procedure detailed in Sadiq et al.\(^ {34} \) with the

| Table 1. The PDB Codes of the Crystal Structures Used to Provide Coordinates for the HIV-1 Protease Bound to Each of the Nine FDA-Approved Protease Inhibitors |
|---------------------------------|-------|-------|
| drug name | abbreviation | PDB |
| Amprenavir | APV | 1HPV |
| Atazanavir | AZV | 2AQU |
| Darunavir | DRV | 2HSI |
| Indinavir | IDV | 1HSG |
| Lopinavir | LPV | 1MUI |
| Nelfinavir | NFV | 1OHK |
| Ritonavir | RTV | 1HXW |
| Saquinavir | SQV | 1FB7 |
| Tipranavir | TPV | 204P |

1230
exception that drug protonation states were determined using Open Babel \(^{37}\) (using a pH of 5, based upon the conditions used in most binding assays for the protease). Gaussian \(^{38}\) was used to perform geometric optimization of the inhibitor with 6-31G** basis functions, and the restrained electrostatic potential (RESP) procedure, also part of the AMBER package, was used to calculate the partial atomic charges. All ligands were found to be neutral except for Indinavir, which was protonated at pH 5, in agreement with the previous findings of Oehme et al. \(^{15}\) (structures for all nine inhibitors are shown in Figure 1). Before the production simulations reported here were run, all systems were minimized and equilibrated for 2 ns using the protocol defined by the BAC. \(^{34}\)

All simulations presented here were performed in the molecular dynamics package NAMD\(^{29}\) in the NPT ensemble with a temperature of 300 K and a pressure of 1 bar, using a 2 fs time step. Free energy was conducted on configuration snapshots generated over the course of MD simulations using the NMODE and MMPBSA modules of the AMBER 9 package. \(^{27}\) Snapshots were output every 10 ps to give 100 snapshots per nanosecond of simulation. Management of ensemble simulation runs and analysis was facilitated by use of the BigJob \(^{40,41}\) extension of the SAGA middleware, \(^{42}\) further details of which are provided in Supporting Information.

The HIV-1 protease consists of two peptide chains, which are usually constructed from the same sequence of amino acids (the structure of the drug bound protein is shown in Figure 2). The catalytic function of the enzyme is performed by a dyad which consists of two aspartic acid (Asp) residues, one in position 25 of each chain (we label that in the first chain D25 and that in the second D25\(^{\prime}\)). It has been established by our group and others that the correct determination of the protonation state of the catalytic dyad of HIV-1 protease is of vital importance in order to obtain accurate binding affinities. \(^{16,43}\) Four possible protonation states are possible for the system: dianionic (D\(^{\text{−}}\)), diprotonated (D25, D25\(^{\prime}\)), Asp 25 protonated (D25), and Asp 25\(^{\prime}\) protonated (D25\(^{\prime}\)). We have employed the same protocol used previously to establish the protonation state appropriate for lopinavir (LPV)\(^{16}\) in order to ascertain the correct protonation state for each drug. Twenty replica simulations (varying only in having different, randomized initial velocities) were performed for each system. Each simulation produced 2 ns of equilibration and 4 ns of production trajectory. In all cases except Tipranavir, where the diprotonated system was favored, the protein was found to be monoprotonated. Full results for each system are given in Table S1.

Once the correct catalytic dyad protonation state was determined for each drug, 30 more replicas were performed for all systems using the same conditions and the binding affinity averages calculated from the full 50 member ensembles used for our final comparison of drug binding affinities. This size of ensemble was shown to provide well converged \(\Delta G_{\text{MMPBSA}}\) results for protease mutants bound to the inhibitor lopinavir in our previous work. \(^{16}\) A second ensemble of the lopinavir system was run and analyzed in order to assess the reproducibility of the
binding affinity estimates produced (the second ensemble is denoted as LPV 2).

3. RESULTS

Our aim in the following sections is to investigate both the level of conformational sampling required to obtain converged results and, once this has been ascertained, to compare the performance of the different methods of free energy calculation in terms of how well they reproduce experimental results.

3.1. Internal Sampling, Convergence, and Reproducibility. The combination of the computational cost of the calculations and the necessity for rapid results in applications such as drug discovery and lead enrichment has led many previous studies of MMPBSA and MMGBSA performance to be limited to single simulations, often providing less than 10 ns of MD trajectory for analysis. Here, we use ensembles of 50 simulations to determine the variability of the results produced by these analyses and how they depend upon the replica length and rate at which conformational samples are taken from the MD trajectories. This strategy is enabled by the increasing availability of supercomputing resources with many thousands of cores (sometimes referred to as petascale resources) which make the execution of many large simulations feasible. In this work, we harnessed resources on both the U.S. XSEDE and E.U. PRACE networks (details of the machines used can be found in the Supporting Information).

3.1.1. Effect of Simulation Length and Sampling Rate. Our previous work indicated that ensembles of 50 replica simulations producing 4 ns of production trajectory were capable of producing well converged MMPBSA values and correctly ranking a series of protease mutants bound to lopinavir (LPV) but that longer single simulations could not. More recent work has suggested that single simulations of 10 ns duration can produce reliable results using MMPBSA for a selection of 14 HIV-1 protease inhibitors selected to give a wide range of affinities and to include representatives of six different scaffolds identified from the literature. Here, we focus on using 50 replica ensembles to see if we can extend our previous analysis to cover all of the FDA approved HIV-1 protease inhibitors. In order to assess the impact of replica simulation length and sampling rate on the convergence of averages obtained by each of the free energy protocols under study, we use ensembles of 50 replicas to see if we can extend our previous analysis to cover all of the FDA approved HIV-1 protease inhibitors. In order to assess the impact of replica simulation length and sampling rate on the convergence of averages obtained by each of the free energy protocols under study, a method to quantify the error in the averages taken from a given ensemble is required. We have chosen to use the statistical technique of bootstrapping. This method involves resampling with replacement the N input data points (in this case, the snapshot values of $\Delta G_{MMPBSA}$, $\Delta G_{MMGBSA}$ or $-T\Delta S_{NM}$) to provide a new bootstrap sample also containing N data points. This process is repeated many times (in our case 10 000 times) and the mean of each bootstrap population calculated. The standard deviation ($\sigma_{boot}$) of these means provides an estimate of the error associated with an average derived from a given sample.

Figure 3 shows the variation of $\sigma_{boot}$ with replica length and sampling rate for $\Delta G_{MMPBSA}$, $\Delta G_{MMGBSA}$ and $-T\Delta S_{NM}$ in the case of the wildtype protease bound to lopinavir (LPV). The most surprising feature of these tracks is that after 4 ns, $\sigma_{boot}$ increases for both $\Delta G_{MMPBSA}$ and $\Delta G_{MMGBSA}$ despite the increased level of sampling being used. This is particularly pronounced for $\Delta G_{MMPBSA}$ and is due to the ingress of water into the active site as identified in our previous work, leading to higher variability in MMPBSA results. This may also explain a slight drift in binding strength observed in the nanosecond average values over time, see Figure S1. The results for $\Delta G_{MMGBSA}$ already appear very well converged at the 4 ns point with a $\sigma_{boot}$ of less than 0.21 kcal mol$^{-1}$ at all of the sampling rates investigated; in the case of $\Delta G_{MMPBSA}$ the error is less than 0.3 kcal mol$^{-1}$ at this trajectory length. We have therefore limited all of the remaining replica lengths to 4 ns. There is little benefit observable for either method when the sampling rate is increased above 20 snapshots per nanosecond. Despite this, for reasons of consistency with our previous work, we have used 100 snapshots per nanosecond for our $\Delta G_{MMPBSA}$ and $\Delta G_{MMGBSA}$ averages. Analysis of the convergence of the separate components of $\Delta G_{MMPBSA}$ and $\Delta G_{MMGBSA}$ can be found in the Supporting Information.

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Figure 3c shows that, unlike for $\Delta G_{\text{MMPBSA}}$, the error on $-T\Delta S_{\text{NM}}$ averages is significantly reduced as the sampling rate increases up to a maximum of 100 snapshots per nanosecond. Here, there is no sign of any change in the convergence behavior as the length of simulation produced by each replica is increased. In order to have comparable error levels at reasonable computational cost, we have chosen to use 25 snapshots per nanosecond for $-T\Delta S_{\text{NM}}$ calculations which provide an estimated error $\sigma_{\text{boot}}$ of approximately 0.3 kcal mol$^{-1}$, comparable to those obtained for $\Delta G_{\text{MMPBSA}}$.

3.1.2. Variance Between Replicas. Given that the vast majority of publications that apply the MMPBSA or MMGBSA methodologies report no more than one or two simulations, it is instructive to examine the variation in binding free energy values that we observe from different replica simulations. Figure 4a,c show the distribution of the averages obtained for each replica via the MMPBSA and MMGBSA methodologies, respectively. The distributions are very broad for all combinations of drug and methodology, in almost every case (excepting NFV using MMGBSA and one of the LPV replicas using MMPBSA) the range of values observed being greater than 7 kcal mol$^{-1}$. Using both methodologies, the values for AZV and RTV have the largest ranges, all of which exceed 10 kcal mol$^{-1}$. In addition to the width of these distributions it is clear from visual inspection that the results do not fit well to the expected Gaussian distribution, which would seem to indicate that sampling on this level is not well converged. By contrast, the $\Delta G_{\text{MMPBSA}}$ and $\Delta G_{\text{MMGBSA}}$ snapshot distributions, shown in Figure 4b and d, respectively, do appear to follow well-defined Gaussian distributions. We suggest that the latter observation and the fact that the distributions for the two LPV replicas lie very close
together mean that the overall averages are well converged, even if this is not clear from the replica distributions.

A similar result is obtained for normal mode estimates of the configurational entropy, albeit with a much larger range in the snapshot values obtained. Again, the snapshot based normalized frequency distributions of the values closely resemble normal distributions with the same mean and standard deviation as shown in Figure 5b. This is a significant result as, previously, we were unable to obtain the expected Gaussian distribution of values when using a lower sampling rate of five snapshots per nanosecond for a total of 1000 snapshots per ensemble (here, we use 5000). The computational cost of normal-mode analysis has meant that many published studies use far fewer snapshots than this. In all cases, the distributions are very broad, the range of values is greater than 80 kcal mol$^{-1}$, and the distributions encompass some snapshots with attractive (negative) contributions to the binding free energy. As for the MMPB(GB)SA results, the replica averages do not produce well-defined normal distributions. However, in line with the bootstrap analysis presented above, the replica distributions are only seen to be as

Figure 5. Normalized frequency distribution analysis of the conformational entropy contribution to the binding affinities ($-T\Delta S_{\text{NM}}$) calculated for (a) each replica and (b) each configurational snapshot computed using normal-mode analysis for each of the nine protease inhibitors studied. In the case of the inhibitor lopinavir (LPV), two replica ensembles are shown, the first with open circles and the latter with filled triangles. Explanations of the abbreviations used for each drug are given in Table 1. The expected normal distribution given the same mean and standard deviation for each data set is shown by the blue lines (the distribution for the second LPV replica is shown in red).

Figure 6. Normalized frequency distribution analysis of the binding affinities calculated for each replica using (a) MMPBSA and normal-mode analysis ($\Delta G_{\text{theor-PB}}$) and (b) MMGBSA and normal-mode analysis ($\Delta G_{\text{theor-GB}}$) for each of the nine protease inhibitors studied. In the case of the inhibitor lopinavir (LPV), two replica ensembles are shown, the first with open circles and the latter filled triangles. The narrowest replica distributions for the $\Delta G_{\text{MMPBSA}}$ and $\Delta G_{\text{MMGBSA}}$ results (for NFV and LPV, respectively) are 1.3 and 1.6 kcal mol$^{-1}$. The expected normal distribution given the same mean and standard deviation for each data set is shown by the blue lines (the distribution for the second LPV replica is shown in red).
broadly distributed as those for the MMPB(GB)SA results. This confirms that by using 25 snapshots per replica we are able to produce well converged estimates of the configurational entropy. Analysis of the three contributions to \(-T \Delta S_{NM}\) indicates that all of the variation observed in the overall computation comes from \(-T \Delta S_{\text{vol}}\) and consequently that the discussion above applies equally to this component of \(\Delta G_{\text{PBAssocNM}}\) and \(\Delta G_{\text{GBAssocNM}}\).

It is not possible to produce consistent per snapshot distributions of \(\Delta G_{\text{theor-PB}}\) or \(\Delta G_{\text{theor-GB}}\) due to the differences in sampling rate, but per replica distributions show similar deviations and have a comparable range to those of \(\Delta G_{\text{MMPBSA}}\) and \(\Delta G_{\text{MMGBSA}}\) (see Figure 6). The narrowest replica distributions for the \(\Delta G_{\text{MMPBSA}}\) and \(\Delta G_{\text{MMGBSA}}\) results (for NFV and LPV, respectively) have a standard deviation of 1.5 and 1.6 kcal mol\(^{-1}\), while those for \(\Delta G_{\text{theor-PB}}\) and \(\Delta G_{\text{theor-GB}}\) (TPV and LPV, respectively) are 2.22 and 2.48 kcal mol\(^{-1}\). The two largest drugs under investigation, RTV and AZV, have consistently broader distributions than the others using either MMPBSA or MMGBSA, with or without the incorporation of \(-T \Delta S_{NM}\).

Unlike MMPB(GB)SA and normal modes, the binding free energy of association, \(\Delta G_{\text{Assoc}}\) has not previously been widely applied, and little is known about the performance of this methodology. Figure 7 shows that we obtain comparatively narrow distributions of the replica averages compared to any of the other free energy components studied here, with a typical range of approximately 5 kcal mol\(^{-1}\). The overall averages also show a relatively small range across the different drugs of only 4.29 kcal mol\(^{-1}\) with six of the nine drugs having averages between 11 and 12 kcal mol\(^{-1}\) (further details are provided in Table S4). It does not appear from the distributions that \(\Delta G_{\text{Assoc}}\) is well converged for the ensembles used here, suggesting that for this technique more sampling is required. An alternative explanation of the poor approximation to a normal distribution is that the naive implementation of this technique applied here (assuming that each replica represents a single local energy minimum) is inappropriate and that some form of clustering might be used before the calculation is performed.

Overall, the results presented here confirm the observation that individual simulation trajectories do not yield reproducible results but that ensembles of simulations can be efficiently used to generate converged free energy estimates. Thus, our findings made previously about lopinavir bound to protease sequences of different binding strengths also apply in the case of the ranking of different protease inhibitors. Remarkably, we observe that the distribution of replica averages of \(-T \Delta S_{NM}\) is no broader than that of \(\Delta G_{\text{MMPBSA}}\) or \(\Delta G_{\text{MMGBSA}}\). Bootstrap analysis of the number of replicas indicates that an ensemble size of 50 represents a good compromise between computational effort and the accuracy of the sampling obtained (see Figure S3).

### 3.2. Comparison with Experimental Data

In order to assess the efficacy of the binding affinity assessment methodologies we have employed, it is necessary to compare our results with experimental values. We searched BindingDB, an online database of measured binding affinities, to identify data sets which covered all nine FDA-approved protease inhibitors under comparable experimental conditions. We found data sets from two groups that fit these criteria, reported by Freire and co-workers (we will refer to this as Expt1) and by Dierynck et al. (we will refer to this as Expt2). Two statistical measures are often used to compare computationally derived binding estimates to those from experiments; the coefficient of determination (\(r^2\)) and the Spearman rank coefficient (\(r_s\)). The former describes the level of linear correlation between the two data sets, the latter whether the rank ordering of the two data sets is the same. Comparing the two experimental data sets, we obtain an \(r^2\) value of 0.47 and an \(r_s\) of 0.90, suggesting that while the rank ordering of the proteins is consistent the exact differences between ligands varies considerably. Consequently, we decided to use \(r_s\) as the main metric to assess the performance of the different free energy methodologies we have employed and to compare our results to both the two experimental data sets and the average of the two (we will refer to the average data set as ExptAvg).

None of the methodologies produce \(r_s\) results higher than 0.63 for Expt1, Expt2, or ExptAvg (see Table 2). The methods incorporating contributions of configurational entropy (\(\Delta G_{\text{theor-PB}}\) and \(\Delta G_{\text{theor-GB}}\)) or the free energy of association (\(\Delta G_{PB\text{-Assoc}}\) and \(\Delta G_{GB\text{-Assoc}}\)) alone generally slightly improve on those that do not (\(\Delta G_{\text{MMPBSA}}\) and \(\Delta G_{\text{MMGBSA}}\)) but with the best results coming from incorporating both (\(\Delta G_{\text{PBAssocNM}}\) and \(\Delta G_{\text{GBAssocNM}}\)). Graphical comparisons of the computed binding

![Figure 7. Normalized frequency distribution analysis of the free energy of association (\(\Delta G_{\text{Assoc}}\)) for each replica. The expected normal distribution given the same mean and standard deviation for each data set is shown by the blue lines (the distribution for the second LPV replica is shown in red).](image-url)

### Table 2. Spearman Rank Coefficient (\(r_s\)) for Each of the Studied Computational Free Energy Methodologies Compared to the Two Experimental Data Sets and Their Average, Ranking All Nine FDA Approved Drugs

| method          | Expt1 | Expt2 | ExptAvg |
|-----------------|-------|-------|---------|
| \(\Delta G_{\text{MMPBSA}}\) | 0.57  | 0.41  | 0.48    |
| \(\Delta G_{\text{MMGBSA}}\) | 0.46  | 0.35  | 0.39    |
| \(\Delta G_{\text{theor-PB}}\) | 0.56  | 0.43  | 0.56    |
| \(\Delta G_{\text{theor-GB}}\) | 0.55  | 0.43  | 0.57    |
| \(\Delta G_{PB\text{-Assoc}}\) | 0.57  | 0.41  | 0.48    |
| \(\Delta G_{GB\text{-Assoc}}\) | 0.51  | 0.38  | 0.44    |
| \(\Delta G_{\text{PBAssocNM}}\) | 0.61  | 0.53  | 0.62    |
| \(\Delta G_{\text{GBAssocNM}}\) | 0.60  | 0.55  | 0.63    |
The binding affinity estimates for two drugs, those for RTV and AZV, stand out as being much too attractive. It was noted above that these two ligands exhibited the least well behaved simulations of smaller ligands. We investigated the idea that simply increasing the length of each replica might overcome any such barriers by extending 10 replicas of the RTV ensemble to 20 ns and found that in one case a conformational flip in the D29 residue allowed water access to the drug which was accompanied by a weakening of \( \Delta G_{\text{MMPBSA}} \) estimates of binding strength of approximately 10 to 15 kcal mol\(^{-1}\) (see Supporting Information). Our previous work has shown that when RTV is docked into protease conformations taken from LPV simulations, a significantly lower binding affinity is obtained than in simulations started from RTV crystal structures\(^{43}\) (although not by enough to result in a correct ranking with respect to the results obtained for other inhibitors in this study). This phenomenon was observed to be linked to the accessibility of the drug and protease active site to individual water molecules. Oehme et al.\(^{35}\) have also suggested that the binding strength of larger drugs is over-estimated by MMPBSA and MMGBSA, although they did not note any specific interactions as the cause.

Excluding the two larger drugs, we obtain much improved rankings as shown in Table 5. Both the \( \Delta G_{\text{FPwAssocNM}} \) and
The recent work of Oehme et al.\textsuperscript{15} examined the ability of MMPBSA and MMGBSA to rank a selection of HIV-1 protease inhibitors (three drugs are common to their work and the present study) using duplicate simulations with similar trajectory lengths to our individual replicas. The range of experimental $\Delta G$ values they attempted to evaluate was approximately double that of the difference between DRV and IDV in our chosen test set. The focus of their work was to investigate the impact of ligand parametrization on the free energy values obtained. They found that the Hartree–Fock method employing 6-31G** basis functions, which we have used in this study (chosen because it was that used to parametrize the rest of the force field we employ), performed particularly poorly; their results produced $r^2$ values with a magnitude of less than 0.01 for $\Delta G_{\text{MMPBSA}}$ and $\Delta G_{\text{MMGBSA}}$. In contrast, their results for $\Delta G_{\text{GBAssocNM}}$ were found to be quite comparable, with $r^2$ values of 0.75 and 0.90 for the Expt1 and Expt2 data sets, respectively.

### Table 5. Spearman Rank Coefficient ($r_s$) for Each of the Studied Computational Free Energy Methodologies Compared to the Two Experimental Data Sets and Their Average

| method                  | Expt1 | Expt2 | ExptAvg |
|-------------------------|-------|-------|---------|
| $\Delta G_{\text{MMPBSA}}$ | 0.80  | 0.71  | 0.72    |
| $\Delta G_{\text{MMGBSA}}$ | 0.63  | 0.63  | 0.60    |
| $\Delta G_{\text{GAssocPB}}$ | 0.73  | 0.78  | 0.82    |
| $\Delta G_{\text{GAssocGB}}$ | 0.67  | 0.74  | 0.76    |
| $\Delta G_{\text{PBAssocNM}}$ | 0.80  | 0.71  | 0.72    |
| $\Delta G_{\text{GBAssocNM}}$ | 0.68  | 0.68  | 0.65    |
| $\Delta G_{\text{GAssocNM}}$ | 0.84  | 0.91  | 0.93    |
| $\Delta G_{\text{GBAssocNM}}$ | 0.75  | 0.90  | 0.87    |

$\Delta G_{\text{GB AssocNM}}$ results provide excellent agreement with both the Expt1 and Expt2 data sets as well as the average results ExptAvg. In all cases, the MMPBSA variant of the analysis method outperforms the MMGBSA alternative. Disappointingly, the inclusion of the free energy of association, $\Delta G_{\text{assoc}}$, alone has no impact on the quality of the ranking using MMPBSA and makes only a slight improvement for MMGBSA.

Considering the more widely used analysis methodologies, the MMPBSA results, including and excluding $-T \Delta S_{\text{MP}}$ provide good agreement with both the Expt1 and Expt2 data sets as well as the average results ExptAvg. MMGBSA without normal modes performs poorly, but the incorporation of $-T \Delta S_{\text{NM}}$ renders its performance comparable to MMPBSA without this contribution. Figure 8 indicates that, when normal modes are not included in the calculation, the relative binding strength for TPV is significantly underestimated compared to the average experimental values for both $\Delta G_{\text{MMPBSA}}$ and $\Delta G_{\text{MMGBSA}}$. The inclusion of the configurational entropy corrects this and allows the tight binding drugs LPV, TPV, and DRV to be differentiated from those which are consistently seen to be weaker binders in the two experimental studies. Furthermore, for this set of ligands, when normal modes are included in the calculation, we significantly improve the coefficient of determination, $r^2$ (see Table 6), indicating that the differentiation between drugs is more quantitatively accurate. The better performance when comparing against Expt2 and ExptAvg compared to Expt1 is largely due to the fact that our results find DRV to be notably more tightly bound than the other drugs, in line with the results of Dierynck et al.\textsuperscript{50} As in the case of $r^2$, the results for $\Delta G_{\text{PB AssocNM}}$ and $\Delta G_{\text{GB AssocNM}}$ outperform the other methods, indicating that there is real value in using $\Delta G_{\text{Assoc}}$ to evaluate the loss of translational and rotational freedom from the simulation trajectory as opposed to using the values produced by normal-mode analysis.

4. CONCLUSIONS

In the present work, we have assessed the potential of the MMPBSA and MMGBSA methodologies to produce the kind of reliable and reproducible predictions of binding affinity that could be used in drug discovery and personalized medicine applications. We have found that for several of the nine FDA-approved HIV-1 protease inhibitors (amprenavir, darunavir, indinavir, lopinavir, nelfinavir, saquinavir, and tipranavir), these methodologies work well, reproducing experimental rankings comparably well to how different experiments correlate with one another.

The two largest ligands in our test set, atazanavir and ritonavir, are poorly handled by all of the approaches we have applied, illustrating that the MMPBSA and MMGBSA methodologies have significant limitations in the range of ligands that can be consistently compared (at least in the case of the HIV-1 protease). Compared to the well ranked ligands, these larger inhibitors have a greater level of interactions beyond the active site. Such interactions distort protein geometry and are poorly handled by the single trajectory approaches which we have employed here. It remains a significant challenge for future work to obtain converged values of the free energy that account for these differences. Our findings also suggest that the optimal sampling strategy and the applicability of the methods will depend strongly on the details of the protein target, with even
what may be thought of as peripheral residues potentially having a significant impact on the computed binding affinities. In order that free energy calculations can be further verified and validated, it is necessary that it becomes standard practice to present an analysis of the sampling and convergence properties of computations, not merely their results. The MMPBSA and MMGBSA methodologies have different convergence properties, with the latter converging much more quickly and being comparatively insensitive to the length of replica simulations in our ensembles. These observations agree with the single simulation results of Srivastava and Sastry, in which greater simulation length was seen to improve the ability of MMPBSA to rank ligands but not MMGBSA. However, we observe that there is greater variability in MMPBSA results as they are extended, which suggests that some of the benefits of trajectory elongation observed in that study may have been fortuitous. Furthermore, 

Figure 8. Average experimental absolute binding free energies ($\Delta G_{\text{ExptAvg}}$) of wildtype HIV-1 protease bound to seven inhibitors compared with theoretical predictions using (a) MMPBSA ($\Delta G_{\text{MMPBSA}}$), (b) MMGBSA ($\Delta G_{\text{MMGBSA}}$), (c) MMPBSA and normal-mode analysis ($\Delta G_{\text{theor-PB}}$), (d) MMGBSA and normal-mode analysis ($\Delta G_{\text{theor-GB}}$), (e) MMPBSA with free energy of association and normal-mode analysis ($\Delta G_{\text{PBAssocNM}}$), and (f) MMGBSA with free energy of association and normal-mode analysis ($\Delta G_{\text{GBAssocNM}}$). RTV and ATZ are excluded due to systematic overestimation of the binding affinity of large ligands. Error bars show the standard errors. The blue line represents a linear regression performed on each data set. Spearman rank coefficients are given in Table 5. A clear improvement in the rankings is produced upon the inclusion of entropy estimates.
Obtaining normal distributions required us to sample 25 conformations per nanosecond of trajectory. Surprisingly, we observe the distribution of average replica free energy contributions from normal modes to be less broad than that for either MMPBSA or MMGBSA.

Within the range of ligands that can be consistently treated, we found that the novel combination of MMPBSA with the vibrational component of normal mode configurational entropy and the free energy of association (as suggested by Swanson et al.\(^{16}\)) provides the best statistical reproduction of experimental results. This methodology captures both the ranking of the drugs (as measured by Spearman rank coefficient, \(r_s\), with a value of 0.93) and the differentiation between drugs (coefficient of determination, \(r^2\), value of 0.92) found in the average results of two experimental data sets. The ranking using MMGBSA alongside these two additional contributions is also superior to other combinations we have evaluated. The inclusion of the computationally inexpensive, but rarely used, free energy of association methodology enhances the ranking obtained considerably, and our results suggest that its calculation should become part of best practice for performing MMPBSA/MMGBSA based calculation in the future.

Good, if notably less accurate, agreement with experimental results was obtained from MMPBSA alone and from both MMPBSA and MMGBSA with normal mode entropy contributions. When the free energy of association is not included in the calculation, the difference produced by including configurational entropy is relatively small. However, the extra computational effort can still be justified by the improvement in coefficient of determination when compared to the average of the two experimental data sets, which suggests that it better reproduces the magnitude of binding affinity difference between the ligands.

One of the most important factors in the ability to obtain the rankings we have presented is the use of ensemble simulations. The values obtained within ensembles of 50 replica simulations of the same protease—drug complex, differing only in initially assigned atom velocities, varied by as much as 10 kcal mol\(^{-1}\), a greater range than that between the best and worst binding FDA-approved inhibitors. The importance of the extensive level of sampling required to obtain consistently converged estimates of binding free energies (irrespective of their accordance with experimental or “real” values) is perhaps the most striking result of the present study. We have presented distributions of the results of all of the free energy computations we have conducted, and we believe that this should become common practice for all future computational binding affinity studies. The reproducibility of the rankings we have obtained is heavily dependent on ensemble size, with a minimum of 25 replicas being suggested by bootstrapping analysis necessary to provide reproducible results. We hope that future work will provide standards that better allow comparisons of different studies using a range of techniques and simulation parameters.

The aim of this study has been not only to validate the ability of MMPB(GB)SA-based free energy calculations to distinguish HIV-1 protease inhibitors but to investigate the broader suitability of the methods for rational drug design and personalized medicine applications. One of the major attractions of MMPB(GB)SA calculations is the promise of general applicability to any chemical species, whereas other techniques, such as free energy perturbation (FEP) or thermodynamic integration (TI), are practically limited to considering free energy differences between molecules that are relatively similar.

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our work demonstrates that, with sufficient sampling, configurational entropies calculated using normal-mode analysis do exhibit normal distributions, contrary to our previous observations.

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Table 6. Coefficient of Determination (\(r^2\)) for Each of the Studied Computational Free Energy Methodologies Compared to the Two Experimental Data Sets and Their Average

| method                | Expt1 | Expt2 | ExptAvg |
|-----------------------|-------|-------|---------|
| \(\Delta G_{\text{MMPBSA}}\) | 0.71  | 0.39  | 0.64    |
| \(\Delta G_{\text{MMGBSA}}\) | 0.48  | 0.26  | 0.44    |
| \(\Delta G_{\text{GB}+\text{PB}}\) | 0.58  | 0.74  | 0.77    |
| \(\Delta G_{\text{GB}+\text{PB}+\text{Assoc}}\) | 0.61  | 0.76  | 0.83    |
| \(\Delta G_{\text{GB}+\text{Assoc}}\) | 0.74  | 0.41  | 0.67    |
| \(\Delta G_{\text{GB}+\text{PB}+\text{Assoc}NM}\) | 0.52  | 0.28  | 0.48    |
| \(\Delta G_{\text{PB}+\text{AssocNM}}\) | 0.66  | 0.86  | 0.92    |
| \(\Delta G_{\text{GB}+\text{AssocNM}}\) | 0.59  | 0.77  | 0.84    |

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Figure 9. The variation of the bootstrap statistics, \(\sigma_{\text{boot-rs}}\), with the number of replicas within an ensemble on the Spearman rank coefficient (\(r_s\)) for the ranking of the seven FDA-approved inhibitors for which our methodologies are valid against the average experimental data set (ExptAvg) using (a) MMPBSA alone (\(\Delta G_{\text{MMPBSA}}\)), (b) MMPBSA and normal-mode analysis (\(\Delta G_{\text{GB}+\text{PB}}\)), and (c) MMPBSA normal-mode analysis and the free energy of association (\(\Delta G_{\text{GB}+\text{Assoc}NM}\)). In all cases, the variation in the ranking is seen to reduce with the number of replicas up to 50, although the decrease slows after 20 to 25 replicas are included in the ensemble.

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In drug discovery and lead enrichment applications, the task is to provide rapid results for libraries of hundreds or thousands of candidate compounds. The computational cost of the ensemble MD simulations that we found necessary to obtain reliable MMPB(GB)SA values means that at present the use of these techniques in this field is unsuitable. It is likely that, for such applications, empirical scoring functions based on quantitative structure–activity relationship (QSAR) models will continue to be the best option (despite the obvious dangers of extrapolating beyond the data used to derive them). In personalized medicine contexts, the demands are however quite different, with normal modes can successfully and reproducibly run concurrently given sufficient computer resources as part of the ViroLab project.17,55 Furthermore, the required turnaround times of days or up to a week are easily feasible. Ensemble simulations represent a very time-efficient method of performing MD calculations, as each replica can be run concurrently given sufficient computer resources. The 6 ns of trajectory for each replica can typically be produced in 15 h (assuming 2.5 h per nanosecond using 64 CPU cores based on the required turnaround times of days or up to a week are easily feasible. Ensemble simulations represent a very time-efficient method of performing MD calculations, as each replica can be run concurrently given sufficient computer resources. The 6 ns of trajectory for each replica can typically be produced in 15 h (assuming 2.5 h per nanosecond using 64 CPU cores based on our experience on several machines) and the free energy analysis (assuming 2.5 h per nanosecond using 64 CPU cores based on trajectory for each replica can typically be produced in 15 h (assuming 2.5 h per nanosecond using 64 CPU cores based on our experience on several machines) and the free energy analysis within another 12 h. These times are only likely to reduce as new technology is introduced and algorithmic improvements are implemented.

Combining the results for different inhibitors presented here with our previous demonstration that the MMPBPSA methodology with normal modes can successfully and reproducibly estimate the binding affinities of mutant proteases to a single inhibitor,16 we believe that MMPB(GB)SA techniques are capable of making significant contributions in academic research as well as basic and clinical medicine.

ASSOCIATED CONTENT

Supporting Information
Details of the SAGA and BigLob middleware used to facilitate this study, alongside thermodynamic decomposition of binding affinities and structural and energetic analysis of the extended ronavir simulations discussed in the main text, are provided. This information 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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