Relative Binding Enthalpies from Molecular Dynamics Simulations Using a Direct Method

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ABSTRACT: The potential for reliably predicting relative binding enthalpies, \( \Delta \Delta E \), from a direct method utilizing molecular dynamics is examined for a system of three phosphotyrosyl peptides binding to a protein receptor, the Src SH2 domain. The binding enthalpies were calculated from the potential energy differences between the bound and the unbound end-states of each peptide from equilibrium simulations in explicit water. The statistical uncertainties in the ensemble-mean energy values from multiple, independent simulations were obtained using a bootstrap method. Simulations were initiated with different starting coordinates as well as different velocities. Statistical uncertainties in \( \Delta \Delta E \) are 2 to 3 kcal/mol based on calculations from 40, 10 ns trajectories for each system (three SH2—peptide complexes or unbound peptides). Uncertainties in relative component energies, comprising solute—solute, solute—solvent and solvent—solvent interactions, are considerably larger. Energy values were estimated from an unweighted ensemble averaging of multiple trajectories with the a priori assumption that all trajectories are equally likely. Distributions in energy—rmsd space indicate that the trajectories sample the same basin and the difference in mean energy values between trajectories is due to sampling of alternative local regions of this superbasin. The direct estimate of relative binding enthalpies is concluded to be a reasonable approach for well-ordered systems with \( \Delta \Delta E \) values greater than \( \sim 3 \) kcal/mol, although the approach would benefit from future work to determine properly distributed starting points that would enable efficient sampling of conformational space using multiple trajectories.

INTRODUCTION

The rapid growth in computational capabilities enables the use of computer simulations to help guide our understanding of biomolecules and their interactions at a level previously unrealized. Estimation of thermodynamic quantities by simulation is particularly important to gain insight into the microscopic details and connect physical interactions with thermodynamic measurements. Here, we consider the direct calculation from molecular dynamics (MD) simulation of the binding enthalpy for a protein—peptide complex. A description of how changes in ligand structure perturb molecular interactions, and hence enthalpy, can provide insight into enthalpy—entropy compensation, or help to explain subtle effects on binding energy when these are difficult to resolve based on crystallographic or NMR structures. 1,2 Furthermore, a strategy to improve the affinity of drug candidates is based on optimizing binding enthalpy and entropy. 3,4 for example, by correlating trends in enthalpy with structural properties such as surface area, chemical composition, etc. This strategy has been challenged on the premise that free energy is more accurately determined experimentally and predicted computationally than enthalpy and entropy, 5 as well as the observation that binding enthalpy is not always predictive of binding free energy. 6,8 Nonetheless, for some systems, knowledge of the binding enthalpy and/or entropy has uncovered information on molecular association that is not present when examining binding free energy alone; distinguishing patterns in enthalpy/entropy can help to understand the molecular properties that affect molecular association. An interesting case is the issue of anticompenensation of entropy and enthalpy in ligand binding, 9,10 to complement the more commonly discussed phenomenon of entropy/enthalpy compensation. In another example, clear trends in binding enthalpy distinguish one group of ligands in a series from another, a trend that is not apparent from free energy alone. 11 Thus, a critical assessment of binding enthalpies can provide insights into the physical factors that govern molecular association, with computational methods involving physics-based models contributing an atomic description of the underlying interactions.

The progress of simulation-based methods to estimate binding free energy is well recognized, 12–19 while the prediction of entropy and enthalpy components remains more challenging. 20,21 The statistical mechanical theory and...
computational methods for the free energy, as well as the decomposition into enthalpic and entropic components of protein–ligand interactions are well described in an insightful review by Levy and Gallicchio. Enthalpy values corresponding to experimental binding measurements can be computed by alternative methods: finite temperature differences to estimate entropy from the temperature derivative of the free energy, and then, the sum of the free energy and entropy times temperature; the derivative methods associated with free-energy perturbation and thermodynamic integration; or a direct estimate from the molecular mechanics energy of end-states. Estimates of the enthalpy and entropy components from derivatives of the free energy function generally are less accurate and have larger errors than estimates of the free energy function itself.

Of the possible approaches to estimate enthalpy, the direct method based on end-states is the most straightforward and offers immediate interpretation of the physical behavior. Nonetheless, the direct method determines the enthalpy of binding from the difference between the energies of the protein–ligand complex and the free molecules obtained from separate simulations. This difference is orders of magnitude smaller than the absolute energy values, and thus, the reliability of the direct method depends on the level of sampling that can be achieved within practical computational times. As such, studies to date using the direct method to estimate protein–ligand-binding enthalpies are few in number to our knowledge, and the alternative method based on finite temperature differences has thus far proven more useful for investigating protein–ligand interactions. A well-designed, seminal study of small-molecule solvation finds that better convergence and more reliable estimates for entropy and enthalpy are achieved with the finite difference method relative to the direct method or derivative quantities. Nevertheless, it should be kept in mind that the accuracy of the finite difference approach is limited by the theoretical assumptions related to the temperature dependence and heat capacity; the finite-difference analysis relates the free energy estimated at different temperatures to the entropy and enthalpy without accounting for changes in the heat capacity, whereas the direct method calculation has the advantage of being carried out at the specified temperature. In addition, the finite difference approach requires that the force field be accurate over the temperature range chosen for the finite difference analysis, which is not generally the case.

Here, we examine the use of the direct method to compute the enthalpy of binding of protein–ligand complexes from individual end-state simulations. A specific estimate of enthalpy is of interest because of the direct comparison with calorimetry data and as the primary factor used to understand structural stability. The direct method has been considered impractical because of the difficulty with convergence of the solvent interactions; however, the increase in computational power suggests that this barrier is rapidly being overcome. Here, the potential for reliably estimating relative enthalpy for protein–ligand binding with the current computer power typically available to academic research groups is considered. The relative enthalpies for three tetrapeptides binding to Src SH2 domain are estimated. Src SH2 is a well-structured 106-residue protein without substantial conformational heterogeneity apparent from NMR heteronuclear relaxation data.

**METHODS**

**Molecular Dynamics Simulations.** The approach to estimate relative binding enthalpies is tested on a model system of three phosphotyrosyl-containing tetrapeptide ligands binding to the 106-residue Src SH2 domain. This association was previously characterized by ITC. The first peptide consists of the canonical Src SH2 recognition sequence (pYEEI). The other two peptides are the conformationally constrained (cpYEEI) or flexible (fpYEEI) mimetics in which the phosphorylated tyrosine residue (pY) is chemically modified (Figure 1A). Equilibrium molecular dynamics simulations were calculated for six systems: three SH2–peptide complexes (SH2–pYEEI, SH2–cpYEEI, SH2–fpYEEI) and three unbound peptides (pYEEI, cpYEEI, fpYEEI).

A set of 40 simulations was generated for each of the six systems. Five sets of starting coordinates for each system were obtained from the crystallographic coordinates of the Src SH2–cpYEEI complex (PDB code 1IS0), containing two copies of the complex in the asymmetric unit, and Src SH2 with a bound 11-residue peptide including the canonical pYEEI sequence (PDB code 1ISPS), containing three copies of the complex in the asymmetric unit. An overlay of the crystallographic structures is shown in Figure 1B. Among these five sets of crystallographic coordinates, the pairwise rms differences between Src SH2 Ca coordinates range from 0.49 to 0.83 Å.

**Figure 1.** Molecules simulated in this study. (A) The three peptides, denoted as canonical (pYEEI), constrained (cpYEEI), and flexible (fpYEEI). (B) Overlay of the Src SH2–peptide complex X-ray structures used to build initial models: 2 chains from the SH2–cpYEEI complex (1IS0, cyan) and 3 chains from the SH2–pYEEI complex (1SPS, magenta). Bound peptides are rendered with thick lines, protein side-chains in thin lines and the protein main-chains in ribbons.
and between all-heavy atoms from 0.97 to 1.54 Å. For the three complexes, the crystallographic ligand was alchemically mutated to the desired pseudopeptide to yield five sets of starting coordinates. Velocities were randomized using random seeds for each of the five starting coordinate sets to establish the 40 unique starting conditions.

SH2 complexes and peptides were prepared for simulations using CHARMM version c35,31 and production runs were performed with NAMD32 using the CHARMM27 all-atom force field33 with CMAP dihedral angle correction.34 Parameters for the nonstandard cpY and fpY residues were described previously.1 Solutes were solvated with 6840 TIP3P water molecules for SH2–ligand complexes, or 2310 water molecules for the free peptides in octahedral boxes, so that the box edges were at least 14 Å from the solute. Nonbonded lists were generated with a 14 Å cutoff and truncation functions applied starting at 10 Å. van der Waals interactions were treated with an atom-based switching function and short-range electrostatics with an atom-based shifting function. Long-range electrostatic interactions were estimated using the particle mesh Ewald method. The energy of the initial systems was minimized for 500 steps with the steepest descent algorithm and then with the adopted basis Newton–Raphson algorithm for 1000 steps or until the energy change between steps was less than 1 kcal/mol, first with harmonic constraints on solute main chain atoms, then finally without constraints.

Molecular dynamics trajectories were calculated with the leapfrog integrator using a 1 fs time step. Constant pressure and temperature (CPT) Nosé–Hoover–Andersen–Klein dynamics used a reference pressure of 1 atm and temperature of 298 K. Simulations were equilibrated for 2500 ps. Production runs of 10 ns were recorded for each of the 40 simulations per system, yielding a total MD run time of 400 ns for each of the six systems. Coordinates were saved every 1 ps. Potential energy values for estimating binding enthalpy were calculated from postprocessing of the trajectories.

Relative Binding Enthalpy Calculation. Relative enthalpies (ΔΔH) for binding various ligands to a single protein were calculated by the direct method of estimating the internal energy of end-states according to Scheme 1. For a protein (P),

$$\Delta\Delta H = \Delta\Delta E = \Delta E_{PL2} - \Delta E_{PL1}$$

where $\Delta E_{PL2}$ and $\Delta E_{PL1}$ are the internal energy of the ligand complexes with the two ligands (L1 and L2), respectively. The total energy of the system is from the molecular mechanics force field of simulations calculated for each protein–ligand complex and the unbound ligand.

The internal energy of the system is from the molecular mechanics force field of simulations calculated for each protein–ligand complex and the unbound ligand. The total system energy, $E_i$, can be partitioned into components corresponding to the geometry and intramolecular nonbonded interactions of the solute, $E_{UV}$, where the solute is either PLn or Ln, the intermolecular interactions between the solute and solvent, $E_{VV}$, and interactions between water molecules, $E_{VV}$:

$$E_i = E_{UV} + E_{VV}$$

where $E_{UV}$ and $E_{VV}$ are the energy of end-states according to Scheme 1. For a protein (P),

$$\Delta\Delta H = \Delta\Delta E = \Delta E_{PL2} - \Delta E_{PL1}$$

with the energy difference, $\Delta E^*$, for each ligand being

$$\Delta E^* = \langle E_{PL1} \rangle - \langle E_{L1} \rangle$$

where $\langle \cdots \rangle$ is the expectation, or time-averaged value obtained from simulations (details given below), and “*” emphasizes the quantity is not the true binding energy given that the energy of the unbound state of the protein $\langle E_{PL1} \rangle$ does not appear in eq 1. An unbound protein simulation is omitted in the analysis because $E_{PL1}$ cancels in the relative binding enthalpies.

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$$E_i = E_{UV} + E_{VV}$$

Mean Energy Values and Uncertainties. A set of trajectories for each state was obtained by calculating multiple, independent MD trajectories. Multiple simulations are considered to provide a broader sampling of conformational space than a single simulation extended for an equivalent total simulation time.35–38 The simulations in this study were calculated independently and not subjected to replica exchange39 or other collective weighting schemes,40 and thus the resulting trajectories serve as a set of repeated measurements of the energy with mean values distributed across the potential energy surface.

The local mean energy, $E_{k,n}$, is estimated from the molecular mechanics force field of the simulations from a single trajectory as the time-average value for trajectory $k$ over the time period corresponding to $N$ snapshots,

$$E_k = \frac{1}{N} \sum_{n=1}^{N} E_{k,n}$$

where $E_{k,n}$ is the energy value of the $n$th snapshot of the $k$th trajectory.

The expected energy value for the set of trajectories, $\langle E \rangle$, is determined from the set of local means, $E_{k,n}$, of the individual trajectories. For $K$ trajectories, the ensemble mean is

$$\langle E \rangle = \frac{1}{K} \sum_{n=1}^{N} E_k = \frac{1}{KN} \sum_{k=1}^{K} \sum_{n=1}^{N} E_{k,n}$$

The uncertainty in the energy values estimated from $K$ independent trajectories was obtained by the bootstrap method41 rather than estimating the error from variances of individual trajectories. For a small sample number, the uncertainty obtained by the bootstrap method is expected to better estimate the width of the underlying Gaussian distribution of mean values than does the standard error. Additionally, applying the bootstrap method to $K$ independent trajectories satisfies the condition of independence among

Scheme 1. Thermodynamic Cycle for the Relative Binding Enthalpy Calculation by the Direct End-State Method

$$P + L_1 \xrightleftharpoons{\Delta E_{n1}}{_{\Delta E_{n2}}} P + L_2$$

$$E_{PL1} \xrightleftharpoons{\Delta E_{n1}}{_{\Delta E_{n2}}} E_{PL2}$$

$$P + L_1 \xrightarrow{\Delta E_{n1}} \xleftarrow{\Delta E_{n2}} PL1$$

$$P + L_2 \xrightarrow{\Delta E_{n1}} \xleftarrow{\Delta E_{n2}} PL2$$

$$\Delta\Delta H = \Delta\Delta E = \Delta E_{PL2} - \Delta E_{PL1}$$

The expected energy value for the set of trajectories, $\langle E \rangle$, is determined from the set of local means, $E_{k,n}$, of the individual trajectories. For $K$ trajectories, the ensemble mean is

$$\langle E \rangle = \frac{1}{K} \sum_{n=1}^{N} E_k = \frac{1}{KN} \sum_{k=1}^{K} \sum_{n=1}^{N} E_{k,n}$$

For the total ensemble here, $K$ equals 40 trajectories and $N$ corresponds to the number of snapshots from a 10 ns trajectory. Certain of the results that follow examine convergence by using cumulative averages for which the value $N$ is varied. Analysis of the simulation ensemble by the simple average defined with eq 5 assumes individual trajectories, $k$, to be equally likely, an assumption discussed in Results.

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\[
\Delta E^* = \langle E_{PL1} \rangle - \langle E_{L1} \rangle
\]
sample observations; thus, the uncertainty of the energy values is reliably estimated using the bootstrap method. Given the set \( A \) of \( K \) local means determined for the independent trajectories, a set \( C \) of synthetic ensemble means, \( \bar{E}_c \), was constructed, where each \( \bar{E}_c \) is the mean of \( K \) values selected randomly from the set \( A \). The uncertainty of the ensemble mean value, \( \delta_{(E)} \), is the standard deviation of the bootstrap sample \( C \),

\[
A = \{ \bar{E}_1, \bar{E}_2, \ldots, \bar{E}_N \}
\]

\[
B_k = \{ x_1, x_2, \ldots, x_k | x_k \in A \}
\]

\[
C = \{ \bar{B}_1, \bar{B}_2, \ldots, \bar{B}_N \}
\]

\[
\delta_{(E)} = \sqrt{\frac{\sum_{k=1}^{N} (\bar{B}_k - \bar{C})^2}{(N - 1)}} \tag{6}
\]

where \( N_c \) is the number of resampled ensembles in the bootstrap sample, and \( \bar{C} \) is the average of the bootstrap sample. In this study, \( N_c \) equals 400. The 95% confidence interval (CI) is calculated from the standard deviation of the bootstrap means: \( 1.96 \delta_{(E)} \). We also calculated the 95% CI from standard error of the local mean values for comparison (see Supporting Information). As expected, the two procedures approach each other as the number of simulations increases so that the bootstrap error and the standard error are similar for our 40 trajectories.

To examine the efficiency for estimating the expected value of the energy, we estimated uncertainty of mean energies for a range of schedules by varying both the number of simulations and the time period of each simulation for a given total simulation time. Bootstrap analysis was applied to randomly selected subsets of trajectories from the ensemble to assess the convergence of the ensemble mean with increasing time per simulation \( (N) \) and number of simulations \( (K) \). We find the mean energy value for \( 2 \leq K \leq 40 \) individual trajectories from 50 to 10,000 ps, or 50 \( \leq N \leq 10,000 \) (eq 4). This set of \( K \) mean values, \( \bar{E}_{k,N} \), is used to generate a bootstrap sample of size 100, and the standard error from sample \( i \), \( \sigma_{(E)} \), calculated by eq 7. The random selection of \( K \) trajectories is repeated 50 times and the final standard error determined from the average variance

\[
\delta_{N,K} = \sqrt{\frac{\sum_{i=1}^{50} (\delta_{(E)}^{(i)})^2}{50}} \tag{8}
\]

The matrix of \( 39 \times 9951 \) standard error values is visualized in a two-dimensional plot.

Two-Dimensional Histograms of Total Energy and rmsd. The potential energy of each system was obtained from postprocessing the corresponding trajectories using the ENER module in CHARMM. For every system, the conformation having the lowest potential energy was chosen as the reference structure for subsequent rms deviation calculations. The rmsd of the backbone heavy atoms (N, Ca, C) with respect to the lowest-energy conformation at every snapshot was calculated using the CORREL module in CHARMM. Then, the ensemble of energy and rmsd values were binned into a two-dimensional histogram to illustrate the distribution of population sampled in the energy–rmsd phase space.

All-Against-All (Pairwise) rmsd Calculations. For every system, the backbone heavy atoms were aligned to the lowest-energy conformation before pairwise rmsd calculations. Weighted pairwise rmsd values were calculated among the 40 aligned trajectories using backbone heavy atoms every 10 ps. Subsequently, calculated pairwise rmsd values were binned into histograms and plotted.

### RESULTS AND DISCUSSION

**Estimates of Relative Binding Enthalpies.** The relative binding enthalpy for three Src SH2–ligand complexes was estimated by the direct method (eq 1) using ensemble-averaging and the a priori assumption that the trajectories sample the same energy basin and have equal likelihood (eq 5). Forty 10 ns trajectories, a total of 400 ns simulation time, were generated from equilibrium molecular dynamics simulations for each of the bound and unbound states of the three Src SH2 ligands illustrated in Figure 1. The two pseudopeptides, cpYEEI and fpYEEI, differ from pYEEI by altering only the pY residue to a constrained isopropyl form or its flexible analogue, respectively. Mean energy values, \( \langle E^3 \rangle \), and the 95% confidence intervals determined from the bootstrap analysis for each system are presented in Table 1, along with the calculated energy differences relative to the Src SH2–pYEEI complex, \( \Delta \Delta E \), and the corresponding experimental enthalpies, \( \Delta \Delta H \).

The 95% confidence interval (CI) from the bootstrap analysis in the mean total energy is \( \pm 1.6 \) to \( \pm 2.6 \) kcal/mol for the complexes, and \( \pm 0.5 \) to \( \pm 0.8 \) kcal/mol for the free peptide simulations. Even though the mean energies for the simulation systems are on the order of \( 10^5 \) kcal/mol, the magnitude of the \( \Delta \Delta E \) values is small (0.14 and 2.2 kcal/mol) and of the same order as the experimental values (1.5 kcal/mol and \( -1.2 \) kcal/mol). Nonetheless, the 95% CI in \( \Delta \Delta E \) is approximately 2 to 3 kcal/mol, which is similar magnitude to the experimental differences, so that rank order cannot be reliably predicted in this case of Src SH2 binding analogue peptides.

| System | Total System Energy \(^{\mathrm{a}}\) (kcal/mol) | \( \langle E^3 \rangle \) | 95% CI |
|--------|---------------------------------|----------------|--------|
| SH2–pYEEI | 73777.8 ± 2.1 | | |
| SH2–cpYEEI | 73741.3 ± 1.6 | | |
| SH2–fpYEEI | 73753.9 ± 2.6 | | |
| pYEEI | 24137.1 ± 0.5 | | |
| cpYEEI | 24100.8 ± 0.6 | | |
| fpYEEI | 24115.5 ± 0.8 | | |

**Table 1. Ensemble Mean Values and 95% Confidence Interval (CI) for the Total Potential Energy, \( \langle E^3 \rangle \), and the Relative Binding Energies, \( \Delta \Delta E \), from Calculations, Along with the Relative Binding Enthalpies, \( \Delta \Delta H \), from Experiment**

| System | \( \Delta \Delta E \) (kcal/mol) | \( \Delta \Delta H \) (kcal/mol) | 95% CI |
|--------|-------------------------------|-------------------------------|--------|
| pYEEI | 0.0 ± 2.2 | | |
| cpYEEI | 0.1 ± 2.2 | | |
| fpYEEI | 2.2 ± 3.2 | | |

**Table 1.**

- **total system energy\(^{\mathrm{a}}\) (kcal/mol)**
- **\( \langle E^3 \rangle \)**
- **95% CI**
- **relative binding enthalpy\(^{\mathrm{b}}\) (kcal/mol)**
- **calcd. \( \Delta \Delta E \)**
- **95% CI**
- **exp. \( \Delta \Delta H \)**
- **\( \delta_{up} \)**

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To better understand the potential for estimating differences in the enthalpies of binding small peptides to a globular protein, in the following subsections we examine the convergence behavior of the mean energy values of the set of trajectories, including the relative binding energies for the SH2–ligand complexes calculated from the MD trajectories. An energy distribution of conformations sampled from multiple trajectories is used to characterize the alternative regions in conformational space visited by the trajectories.

Convergence and Certainty of Energy Values and Relative Binding Energy. The direct method for estimating the relative enthalpy of binding, \( \Delta \Delta E = \Delta E_{\text{complex}}^* - \Delta E_{\text{peptide}}^* \), depends on the total potential energy of the system for the SH2–peptide complexes and free peptide ligands (Table 1). Thus, the ability to resolve the relatively small energy differences of \( \Delta \Delta E \) from the large \( E^T \) values relies on determining the average values of the energies with high confidence. A necessary condition for this reliability is that the value be well converged, although it is recognized that such convergence does not guarantee sampling is complete in a true global sense and can be an indicator of only a localized sampling. The ensemble-averaged value as a function of the production time per simulation used for averaging is one indicator for the convergence of the estimated value. This cumulative ensemble average for \( \Delta \Delta E_{\text{ref}} \) as well as for \( E^T \) of the complexes and peptides and \( \Delta E_{\text{ref}}^* \), is shown in Figure 2. The limiting slopes (Table 2) for \( E^T \) are approximately \(-0.2\) to \(-0.6\) kcal/mol/ns for the complexes and \(0.02\) to \(0.08\) kcal/mol/ns for unbound peptides. These cumulative averages of the energy from the ensemble mean over 40 trajectories are considerably more stable than the cumulative average of a single trajectory; single-trajectory cumulative averages fluctuate over much larger magnitudes and have limiting slopes that range from \(2.1\) to \(-2.3\) kcal/mol (see Supporting Information). The absolute difference in the ensemble-mean energy, \( \Delta E_{\text{ref}}^* \), Figure 2C continues to drift at the end of the 10 ns simulation time period as a result of the difference in the limiting slopes for the complex and peptide ensemble averages. The drift in \( \Delta E_{\text{ref}}^* \) partially cancels when comparing two complexes so that the relative energies, \( \Delta \Delta E \), plotted in Figure 2D with pYEEI as the reference ligand 1, appear better converged with flatter curves at shorter times. Nevertheless, smaller limiting slopes in \( E^T \) over a longer simulation time period would yield greater confidence in the calculated values for \( \Delta \Delta E \) and estimating differences of one to two kcal/mol. The limiting slopes in Figure 2A suggest that 400 ns molecular dynamics simulations are not sufficient to exhaustively sample the complex superbasin of the potential energy surface of these reasonably “simple” protein complexes.

The dependence of the statistical certainty of the energy estimates on simulation time was determined using the bootstrap method41 (see Methods). The decrease of the 95% CI for \( \langle E^T \rangle \) is shown in Figure 3 for the Src SH2–pYEEI with a 2-dimensional plot as a function of the length of the individual trajectories and as a function of the number of trajectories, for which a subset of the 40 trajectories is used in the bootstrap analysis. Black contour lines in the figure denote a constant computer simulation time according to the combined number and length of the individual trajectories. Analogous plots are provided in Supporting Information for the other systems. The uncertainty converges approximately as expected from sampling a single statistical population; the 95% CI for 100 ns computer time is ranges from \(3\) to \(4\) kcal/mol and for 400 ns computer time is \(2.2\) kcal/mol. The efficiency for diminishing the statistical uncertainty is nearly uniform along the contour lines for computer times greater than 200 ns, so that increasing the

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**Figure 2.** Convergence of the 40-trajectory ensemble averages (eq 5) accumulated over increasing production time of the simulations for various energies: (A) the total energy (\( E^T \)) for the complexes, (B) total energy for the ligands, (C) effective binding enthalpies (\( \Delta E_{\text{ref}}^* \)), and (D) relative binding enthalpies (\( \Delta \Delta E \)). Red, SH2–pYEEI or pYEEI; blue, SH2–cpYEEI or cpYEEI; green, SH2–fpYEEI or fpYEEI. Points in C are shifted by subtracting the final value of \( \Delta E_{\text{ref}}^* \) for pYEEI (10 ns/simulation).

**Table 2.** Least-Squares Fitted Slopes at the Long-Time Limit of the Ensemble Mean Cumulative Averages, the 8 to 10 ns Regions in Figure 2

|                  | \( \langle E^T \rangle \) | \( \langle \Delta E^T \rangle \) | \( \langle \Delta \Delta E^T \rangle \) |
|------------------|---------------------------|-------------------------------|--------------------------------------|
| pYEEI            | -0.16                     | -0.26                         | 0.000                                |
| cpYEEI           | -0.56                     | -0.56                         | 0.44                                 |
| fpYEEI           | -0.32                     | -0.32                         | -0.16                                |
time period of the simulation or the number of simulations is equally effective.

In addition to the convergence of $E^T$, the convergence of the estimate for the components of the total potential energy was evaluated. The cumulative ensemble-averaged values for the difference in binding enthalpy contributed by solute–solute ($E^{UV}$), solute–solvent ($E^{VU}$), and solvent–solute ($E^{UV}$) interactions are shown in Figure 4. (Note the large scale of A relative to the panels B–D). Because the total energy is constant but partitioned among these three components, the convergence is slower depending on the relaxation time for partitioning. The convergence behavior manifests in the time dependence of the cumulative average of the ensemble component energies is therefore poor and the estimates vary tens of kCal/mol over the full 400 ns simulation time.

**Overlap of Trajectories in Conformational Space.** The cumulative ($E^T$) values of individual trajectories vary, as noted above and shown in Supporting Information. This behavior raises the question of whether the trajectories sample the same energy basin in the conformational space of the bound complex, as assumed a priori for ensemble averaging. To gain insight into the actual overlap and nature of the conformational regions populated by individual trajectories, we examine distributions in an energy–rmsd system, a natural choice to address this question. The energy is the total potential energy and the rmsd value is the pairwise root-mean-square difference in the N, Ca, C atoms of the SH2 backbone between the trajectory snapshots and the structure with the lowest potential energy in the 40-trajectory ensemble.

Presented in Figure 5 are the projected distributions in the energy–rmsd space for the three Src SH2–peptide complexes. Results for one complex are shown in a group of two rows: SH2–pYEEI is the top group, SH2–cpYEEI is the middle group, and SH2–fpYEEI is the bottom group. Each plot in the top row displays the projection for simulations initiated from five different sets of coordinates, while the plots in the bottom row correspond to the same individual simulations but separated by those initiated from one set of coordinate with five different velocities. (Twenty-five of the forty trajectories are shown to facilitate comparison of the initial conditions that vary by either coordinates or velocities.) Thus, a distribution in the top rows indicates the coverage of conformational space achieved by starting simulations with different initial coordinates, while that in the bottom rows reflects the coverage achieved by starting simulations with one initial coordinate set and varying velocities, as is more typically done to generate multiple trajectories. The fluctuations in energy values cover a broad range of roughly 500 kcal/mol for values of rmsd that mostly vary in the region from 1.0 to 1.5 Å. Some distributions in panel A through J show one peak with a maximum that is more or less centrally positioned, while others show two peaks distinguished along the rmsd coordinate. One might anticipate that multiple trajectories initiated with different initial coordinates (top rows A–E) might show greater dispersion in the energy–rmsd distribution than multiple trajectories initiated with the same coordinates but varying velocities (bottom panels F–J); however, that is not the case: the dispersions in the distributions do not differ significantly.

Nearly all of the individual trajectories visit two or more regions in energy–rmsd space (see Supporting Information),
consistent with different trajectories sampling overlapped parts of conformational space and that the barriers in the underlying potential energy surface giving rise to differences between the trajectories are low. In addition, the combined population from the 40-trajectory ensemble (panel K) is one broad peak in energy−rmsd space with a single maximum indicating the overlap of individual trajectories. Together, the energy−rmsd distributions are consistent with the individual trajectories sampling one superbasin in conformational space, albeit sampling different parts of that basin, and thus support this assumption for ensemble averaging.

The different regions apparent in plots A−J are close in energy for a given Src SH2−peptide complex. For each of the three complexes, regions in energy−rmsd were determined from the distinct peaks in panels A−J, and the mean energy calculated from the conformations in the 400 ns ensemble falling into a given region. The average energies and populations for each energy−rmsd region are listed in Table Figure 5.
3. The mean energies differ by a few kcal/mol; the largest difference is approximately 7 kcal/mol. Thus, the peak regions in Figure 5 are nearly degenerate in energy.

| Table 3. Average Energy of Snapshots Falling in the Different Peak Regions Populated in the Energy—rmsd Space of the Src SH2 Complexes |
|---------------------------------------------------------------|
| complexes | rmsd range (Å) | \( E^T \) (kcal/mol) | \( \sigma^k \) (kcal/mol) | population<sup>a</sup> |
|-----------------|----------------|-----------------|----------------|------------------|
| SH2–pYEEI       | 0.42 to 1.25   | −73779.0        | 132.0          | 231 461          |
|                 | 1.25 to 1.96   | −73776.2        | 132.4          | 168 539          |
| SH2–cPVEE1      | 0.46 to 1.08   | −73742.0        | 131.9          | 93 600           |
|                 | 1.08 to 1.58   | −73741.2        | 132.4          | 281 492          |
|                 | 1.58 to 2.32   | −73740.2        | 132.4          | 24 908           |
| SH2–fPVEE1      | 0.45 to 1.08   | −73756.6        | 132.8          | 170 977          |
|                 | 1.08 to 1.38   | −73752.7        | 132.5          | 164 773          |
|                 | 1.38 to 1.83   | −73750.0        | 132.5          | 52 797           |
|                 | 1.83 to 2.50   | −73749.2        | 133.1          | 11 453           |

<sup>a</sup>The rmsd range is estimated for the peak regions. \( \sigma^k \) is the energy standard deviation. \( ^a \)Number of snapshots in the peak region.

## CONCLUSION

The relative enthalpies for the binding of the 106-residue Src SH2 domain to three flexible peptides were estimated using the direct method of MD determined from the end-states. Notably, the estimated \( \Delta \Delta E \) values were of similar magnitude as the differences in the experimental enthalpies, which are only 1–3 kcal/mol. Nonetheless, the rank order was not predicted correctly for these three complexes. The statistical error in the estimates for \( \Delta \Delta E \) from 40 10 ns simulations for each bound and unbound peptide system is 2 to 3 kcal/mol (Table 1). Based on cumulative ensemble-averaged values (Figure 2), the end-state energies are not fully converged; the cumulative averages for \( E^T \) of the complexes have a limiting rate of change from 0.2 to 0.6 kcal/mol-ns, although there is some cancellation in the \( \Delta \Delta E \) values so that these appear better converged. These results support the application of the direct method of MD simulations to predict relative binding enthalpies for \( \Delta \Delta H \) values greater than 3 kcal/mol. Longer simulations are expected to yield improvements over the predictions reported here, which correspond to 400 ns total simulation time. Likely, the most successful application of the direct method of MD simulations for predicting \( \Delta \Delta H \) values will be to complexes that are conformationally well-ordered and absent of longer-time scale conformational fluctuations that substantially alter the protein energetics.

Multiple trajectories starting from closely related but alternative configurations, or different velocities exhibit different \( E^T \) values with poorer convergence than the ensemble-averaged values. That individual trajectories have different energies suggests that trajectories need to be weighted.35 How to weight trajectories is an important but difficult consideration. The analysis here has utilized a simplistic, unweighted ensemble averaging with the a priori assumption that all trajectories are equally likely. Clustering in energy—rmsd space indicates the trajectories sample the same basin and the apparent difference in mean energy value between trajectories is due to sampling alternative local regions of this superbasin. Clearly, the sampling is incomplete. For more reliable estimates of binding enthalpies, additional theoretical development is needed; in particular, the direct method would gain from reliable methods to achieve canonically distributed starting conditions both in terms of reaching statistical certainty efficiently and, importantly, complete sampling for accurate ensemble-averaged end-state energy values.

An advantage of the direct method for estimating the relative binding enthalpies is its basis on the total energy of the system rather than a summation of various energy terms, for example solvation energy plus protein internal energy. The fluctuations in the total energy are small relative to the fluctuations of any set of component energy terms; component terms freely exchange energy and thus exhibit large fluctuations. Therefore, the certainty in the total energy converges more rapidly while component energies by nature are more difficult to converge. Nonetheless, decomposition of thermodynamic values can provide useful insights into whether quantification is not essential. An example we note here is the observation that the fluctuations in the cumulative mean values for the components \( \Delta \Delta E^{UV} \), \( \Delta \Delta E^{VV} \), and \( \Delta \Delta E^{UVV} \) have the same convergence behavior and are close to being perfectly correlated either negatively (\( \Delta \Delta E^{UV} \) with either \( \Delta \Delta E^{UU} \) or \( \Delta \Delta E^{VV} \)) or positively (\( \Delta \Delta E^{UU} \) with \( \Delta \Delta E^{VV} \)) (Figure 4). Correspondingly, an inverse correlation of \( E^T \) and \( E^U \) is observed upon examination of the mean values from individual trajectories (Supporting Information). The \( E^U \) variance among the multiple trajectories is large compared to \( E^T \) and approximately equal to the sum of the variances in \( E^T \) and \( E^U \). This macroscopic behavior is reminiscent of the microscopic property whereby the reaction field solvation energy cancels the Coulombic interaction.

## ASSOCIATED CONTENT

Supporting Information

Plots of the cumulative average of \( E^T \) for the individual trajectories of the six simulations systems (Figure S1); plots analogous to Figure 4 showing 95% CI results for the ensemble-averaged \( E^T \) values for SH2–cPVEE1, SH2–fPVEE1 (Figure S2), and the three unbound peptide ligands (Figure S3), as well as the ensemble-averaged \( E^{UU} \), \( E^{VV} \), \( E^{UUV} \), values for the three complexes (Figure S4) and unbound peptide ligands (Figure S5); plots of pairwise, or all-against-all, rmsd distributions to examine overlap of individual trajectories in sampling conformational space (Figure S6); a plot showing the deviation from the ensemble mean energy of the local mean energy from individual trajectories, where the energy is the total potential, \( E^T \), or the highly correlated component terms \( E^{UU} \), \( E^{VV} \), and \( E^{UUV} \) (Figure S7); a plot showing 95% CI from standard error calculation for ET values (Figure S8) and for \( E^{UU} \), \( E^{VV} \), \( E^{UUV} \) values (Figure S9) for complex SH2-pYEE1. Table S1 lists the limiting slopes in the cumulative potential energy of forty trajectories for each of the three SH2-peptide simulations. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

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