Multidimensional Potentials of Mean Force from Biased Experiments Along a Single Coordinate

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Abstract

External biasing forces are often applied to enhance sampling in regions of phase space which would otherwise be rarely observed. While the typical goal of these experiments is to calculate the potential of mean force (PMF) along the biasing coordinate, here I present a method to construct PMFs in multiple dimensions and along arbitrary alternative degrees of freedom. A protocol for multidimensional PMF reconstruction from nonequilibrium single-molecule pulling experiments is introduced and tested on a series of two dimensional potential surfaces with varying levels of correlation. Reconstruction accuracy and convergence from several methods - this new protocol, equilibrium umbrella sampling, and free diffusion - are compared, and nonequilibrium pulling is found to be the most efficient. To facilitate the use of this method, the source code for this analysis is made freely available.
1 Introduction

External biasing forces are often applied to enhance sampling in regions of phase space which would otherwise be rarely observed.¹ These perturbing forces are applied in both time-dependent, as in single-molecule pulling²–⁵ and steered molecular dynamics⁶,⁷ experiments, or time-independent forms, as in umbrella sampling simulations.⁸ Usually, the direction of bias is chosen to follow the reaction coordinate for an interesting process, such as a large conformational change in a biological macromolecule. With judicious reweighing of biased ensemble properties, previous workers have reconstructed the potential of mean force (PMF) along this bias coordinate.⁹–¹⁵

The ability to reconstruct PMFs in multiple dimensions and along multiple alternative degrees of freedom would significantly expand the data analysis tool kit for biased experiments. Although the appropriate choice of reaction coordinate is fundamental to kinetic calculations, transition state theory, and the accurate measurement of biased data is collected along an initial, possibly arbitrary choice. In addition, by using multidimensional PMFs, the mutual entropy¹⁶ can be used quantify and compare correlations between modes. Here, I develop methods to perform these calculations. They are applicable to both computer simulations, in which system properties are completely known, or laboratory experiments in which several properties can be simultaneously measured.

2 The Potential of Mean Force

The PMF, $F_0(x)$, along a single coordinate, $x$, is defined as

$$e^{-\beta[F_0(x)+\delta F]} = \frac{\int \delta(x-x(r))e^{-\beta[H_0(r)]} \, dr}{\int e^{-\beta[H_0(r')]} \, dr'}$$

where $\delta F$ is an arbitrary constant, $\beta = 1/k_B T$, and $H_0(r)$ is the unperturbed system energy as a function of the phase space position $r$.

In two dimensions, $F_0(x)$ becomes $F_0(x, y)$ and $\delta(x-x(r))$ is replaced by $\delta(x-x(r))\delta(y-y(r))$. Generalizing to multiple dimensions, $F_0(x)$ becomes $F_0(x, y_1, y_2, ..., y_n)$ and the delta function is replaced by $\delta(x-x(r))\prod_{i=1}^{n} \delta(y_i - y_i(r))$, where $n$ is the number of dimensions. If the PMF encompasses all system dimensions, it can simply be referred to as the potential.

3 Multidimensional PMFs from Nonequilibrium Experiments

Multidimensional PMFs can be reconstructed from time-dependent biasing experiments by applying the nonequilibrium work relation¹⁷,¹⁸ in a manner analogous to Hummer and Szabo.¹¹,¹² In a single-molecule pulling experiment with the Hamiltonian $H(r, t) = H_0(r) + V[r, t]$, where $V[r, t]$ is the time-dependent external bias, the biased probability density function is

$$e^{-\beta[H_0(r)+V[r,t]]} \int e^{-\beta[H_0[r']]+V[r',t]} \, dr' = \langle \delta(r-r(t))e^{-\beta W_t} \rangle$$

where $W_t$ is the accumulated work, $\int_0^t \frac{\partial H(r,t)}{\partial r} \, dt$.¹¹,¹²

To calculate the PMF along the pulling coordinate, Hummer and Szabo consider the special case that the perturbation depends only on the biasing coordinate and time, $V(r, t) = V[x(r), t]$. They then multiply equation 2 by $e^{\beta V(x,t)}\delta(x-x(r))$, integrate over all phase space, and use the definition in equation 1 to obtain

$$e^{-\beta F_0(x)} = \langle \delta(x-x(r(t)))e^{-\beta(W_t-V[x,t])} \rangle$$
If we instead multiply equation\textsuperscript{[2]} by a two-dimensional delta function, we then obtain the two-dimensional PMF. Similarly, multiplying by a multidimensional delta function will lead to a multidimensional PMF. (Although this is possible in principle, high dimensionality will lead to greater difficulty with sampling.) In the two-dimensional case, the PMF is

\begin{equation}
 F_\alpha(x, y) = -\beta^{-1} \ln \left( \frac{\sum_t \langle \delta(\mathbf{x} - \mathbf{x}(t)) \delta(\mathbf{y} - \mathbf{y}(t)) e^{-\beta W_t} \rangle}{\sum_t \langle e^{-\beta W_t} \rangle} \right)
\end{equation}

In any particular time slice, the position will be more highly sampled near the minima of the bias potential. To improve accuracy over the entire range of \(x\), data from different time slices can be combined by adapting the weighted histogram analysis method (WHAM).\textsuperscript{[9]} After unbiasing with respect to the accumulated work as well as the time-dependent potential,\textsuperscript{[11,12]} the resultant PMF is

\begin{equation}
 U(x, y, \alpha) = \alpha U_a + (1 - \alpha) U_b
\end{equation}

where \(U\) is in units of \textit{pN nm} and positions \(x\) and \(y\) are in units of \textit{nm}. \(U_a\) and \(U_b\) both have minima at (0,0) and (45,30), and \(U_a\) has additional minima at (0,30) and (45,0). Part \textit{c} of figures\textsuperscript{[11,12,3]} and\textsuperscript{[4]} show the potential \(U(x, y, \alpha)\) at \(\alpha\) of 0, 0.30, 0.75, and 1.00, respectively.

In \(U_a\) (Fig.\textsuperscript{[11]}), \(x\) and \(y\) are completely independent, while in \(U_b\) (Fig.\textsuperscript{[12]}), they are highly correlated. This is quantified by the mutual entropy (Fig.\textsuperscript{[5]}), which was calculated by assuming a Boltzmann distribution for \(x\) and \(y\), \(f(x, y) = e^{-\beta U(x, y)}\), and numerically integrating equation\textsuperscript{[6]} between \(-20 < x < 65\) and \(-20 < y < 50\) via adaptive Simpson quadrature.

As expected, mutual entropy decreases as the potential becomes more like \(U_a\). The joint entropy, \(S_{xy} = \int f(x, y) \, dx \, dy\), and the marginal entropies, \(S_x = \int f(x) \, dx\) and \(S_y = \int f(y) \, dy\), are mostly stable as a function of \(\alpha\). These magnitude of these factors cannot be directly compared to the mutual entropy because they are offset by an arbitrary constant, whereas mutual entropy is invariant to scale.
Figure 1: PMF reconstruction on the surface with $\alpha = 0$ from umbrella sampling. 

- **a.** PMFs along the $y$ coordinate. The blue line is the theoretical surface, the red asterisks are the biased reconstruction, and green dots are the naïve reconstruction.
- **b.** Contour plot of the reconstructed potential.
- **c.** Contour plot of the theoretical potential. The scale for both b and c is indicated by the side color bar.
- **d.** PMFs along the $x$ coordinate. Annotation is the same as in **a.**

Figure 2: PMF reconstruction on the surface with $\alpha = 0.3$ from time-dependent biasing experiments. Annotation is the same as in Figure 1.
Figure 3: PMF reconstruction on the surface with $\alpha = 0.75$ from diffusion experiments. Annotation is the same as in Figure 1 except there are no WHAM reconstructions.

Figure 4: PMF reconstruction on the surface with $\alpha = 1.0$ from time-dependent biasing experiments. Annotation is the same as in Figure 1.

Figure 5: Entropy as a function of the coupling parameter $\alpha$. The mutual entropy, $S_{\text{mutual}}$, is designated by connected squares. The joint entropy, $S_{xy}$, and the marginal entropies, $S_x$ and $S_y$, are labeled and marked by connected circles.
error and mutual entropy (Fig. 6), as other factors of the free energy surface come into play. There is no simple dependence, however, between unbiased diffusion on the surface are indicated by diamonds. Errors in WHAM-reconstructed surfaces are connected by solid lines, while the naïve estimate errors are not connected.

5 Simulations on a Two-Dimensional Energy Surface

Brownian dynamics simulations were run on the series of potential surfaces. The progress of a dynamical variable is given by $\Delta x(t) = \frac{F(x(t-1) + D\Delta t)}{k_B T} + (2D\Delta t)^{1/2}R$, where $F(x) = -\frac{\partial U(x,y)}{\partial x}$ is the force, $T$ is the temperature (300 K), $D$ is the diffusion constant (1200 nm$^2$/s), $\Delta t$ is the time step (1 ms), and $R$ is a normally distributed random variable.

After 1000 steps of equilibration, umbrella sampling and pulling simulations were run for 100 iterations of 10000 time steps. In addition, 10 unbiased diffusion simulations were run for 100000 time steps. Umbrella sampling was performed with a harmonic bias, $V(x) = 1/2k_s(b - x)^2$, setting the spring constant, $k_s$, at 0.1pN/nm and the bias center, $b$, at one of 100 evenly spaced positions between -5 and 50 nm. Pulling simulations were performed with the time-dependent perturbation, $V[x(r), t] = 1/2k_s(b(t) - x(r))^2$, with $k_s$ set to 0.4pN/nm. A periodic biasing program, $b(t) = 22.5 - 27.5 \cos(2\pi t/10000)$, which has one period per trajectory, was used. Accumulated work was numerically integrated by $W_t = \sum_{j=1}^{t} -k_s[(x_j + x_{j-1})/2 - (b_j + b_{j-1})/2](b_j - b_{j-1})$, where $b_j$ is the position of the bias center at time step $j$.

Theoretical surfaces, $U(x)$ and $U(y)$, were calculated by assuming a Boltzmann distribution, $f(x,y) = e^{-\beta U(x,y)}$, and numerically integrating $f(x,y)$ over the other coordinate (with limits of $-20 < x < 65$ and $-20 < y < 50$ using adaptive Simpson quadrature to obtain a marginal probability density. PMFs are then obtained by taking natural logarithms and multiplying by $-k_B T$. In time-dependent biasing experiments, reconstructed surfaces for $x$ were calculated using equation 3 two-dimensional surfaces from equation 5 and surfaces along $y$ by integrating over $x$; from the estimate of $f(x,y)$, the sum is taken over $x$ to yield an unnormalized $f(y)$, from which $U(y) = -k_B T \ln f(y)$ is calculated. PMFs from umbrella sampling were calculated as described by Roux. Minima of all PMFs were set to zero. Errors were calculated by the root mean square deviation from the theoretical surface at 50 points between $-10.9091 < x < 55.9091$ and $7.2727 < y < 37.2727$, ranges which include the zeroes of $U_a$ and $U_b$. The root mean square error (RMSE) along $y$ is defined as $\sqrt{\frac{1}{50} \sum_{i=1}^{50} [F_{\text{theoretical}}(y_i) - F_{\text{reconstructed}}(y_i)]^2}$.

6 Results and Discussion

When $x$ and $y$ are completely independent, the naïve estimate for $F(y)$ is as good as the estimate from equation 5 (Fig. 4a). If $x$ and $y$ are highly correlated, however, the naïve estimate fails and a WHAM-informed reconstruction method is necessary (Fig. 1). There is no simple dependence, however, between error and mutual entropy (Fig. 6), as other factors of the free energy surface come into play.
Figure 7: Root mean square error for $y$ PMF estimates as a function of number of simulations, $n$. Error bars indicate the mean and standard deviations of RMSE from five reconstructions using $n$ simulations. Reconstruction errors for pulling experiments are indicated by squares and for umbrella sampling by circles. Unbiased simulations were run for 10 times as long and are therefore spaced 10 times farther apart on the x axis.

For example, due to the highly diffusive nature of the dynamics and the large timestep necessary for fast sampling, sampling is erroneously improved near the boundaries of small low-energy wells. This leads to systematic free energy underestimates in all simulations, even those without any external bias (i.e. Fig. 3). Although there are errors in WHAM-informed reconstructions from simulations with external bias, they are no worse than errors from long simulations with free diffusion (Fig. 5).

On certain free energy surfaces, phase space sampling along the $y$ dimension may be limited by trapping in local minima. This is especially true if $x$ and $y$ are relatively independent and biasing along $x$ does not enhance conformational sampling along $y$. Sampling is generally limited in high-energy regions of phase space (see Figs. 1, 2, 3, and 4). In these regions, the most accurate two-dimensional PMF reconstruction would require biasing along both $x$ and $y$, which can be accounted for by a simple modification of equation 5. In general, accurate PMF reconstruction in higher dimensions may require biasing along coordinates that closely correspond to these dimensions. Close correspondence may be indicated by high mutual entropy; this aspect of analysis will be left to further study.

On the toy two-dimensional surfaces studied, PMF reconstructions along $y$ using equation 5 actually converge more quickly than equilibrium umbrella sampling and reconstruction from unbiased experiments (Fig. 7). While time-dependent biasing experiments will sample the whole span of the bias coordinate de facto, umbrella sampling will inherently require a certain number of simulations at different bias centers to attain this broad range. Similarly, unbiased simulations will need to be run for longer in order to reach more distal regions of phase space. Generally, it is preferable to run a greater number of nonequilibrium experiments than a single long equilibrium experiment because the outcome of each individual trajectory can be highly dependent on initial conditions. These sampling problems described on two-dimensional surfaces are likely to be enhanced in systems with higher complexity. The main caveat with this method, as with all methods based on the nonequilibrium work relation, is that with greater free energy differences, the likelihood of observing a trajectories with negative dissipative work (work that is less than the free energy difference) is increasingly unlikely.

Pulling experiments are poised to find broader utility in studies of nanoscale systems, particularly when free energy differences between states are not prohibitively large. I hope this work will find wide application in computational and laboratory experiments and stimulate further research in the area. MATLAB source code to perform the described analysis is available at http://mccammon.ucsd.edu/~dminh/software/.
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