Fast Adaptive Flat-histogram Ensemble for Calculating Density of States and Enhanced Sampling in Large Systems

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We presented an efficient algorithm, fast adaptive flat-histogram ensemble (FAFE), to estimate the density of states (DOS) and to enhance sampling in large systems. FAFE calculates the means of an arbitrary variable \(U\) in generalized ensembles to form points on the curve \(\beta_s(U) \equiv \frac{\partial S(U)}{\partial U}\), the derivative of the logarithmic DOS. Unlike the popular Wang-Landau-like (WLL) methods, FAFE satisfies the detailed-balance condition throughout the simulation and automatically generates non-uniform \((\beta, U)\) data points to follow the real change rate of \(\beta_s(U)\) in different \(U\) regions and in different systems. Combined with a \(U\)-compression transformation, FAFE reduces the required simulation steps from \(O(N^{1/2})\) in WLL to \(O(N^{1/2})\), where \(N\) is the system size. We demonstrate the efficiency of FAFE in Lennard-Jones liquids with several \(N\) values. More importantly, we show its abilities in finding and identifying different macroscopic states including meta-stable states in phase co-existing regions.

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\[ \text{where } r_k \text{ is the } k^{th} \text{ conformation of the sample } [13], \text{ and } w_k \text{ is the weight factor defined as: } w_k = e^\left[ S(U(r_k)) - f(X(r_k)) \right]. \] The statistical error of this weighted mean is proportional to \(B^{-1/2}\), where \(B\) is the effective size of the weighted sample, \(B = \sum w_k^2 / \sum w_k^2\). Only when sufficient conformations of the sample are located inside the important regions of the \(f(X)\) ensemble (i.e., \(B\) is large enough), will the weighted sample provide a good estimate of the ensemble mean.

The Wang-Laudau (WL) algorithm \(8\) estimates \(S(U)\) by updating \(\tilde{S}(U)\) from an initial guess until the visited histogram is roughly a constant. The WL algorithm can be easily implemented and has been widely applied \(10, 11, 12\). However, two features of this algorithm would limit its applicability. First, the required number of simulation steps is at least \(O(N^2)\), where \(N\) is the size of system; hence simulations of large systems will be costly. This estimate arise from the fact that the area below the curve \(S(U)\) is order-\(N^2\) (because both \(S\) and \(U\) are proportional to \(N\)), which must be filled by using the small kernel functions with presumed width and height in WL \(3\). Second, detailed balance is not satisfied in the WL algorithm \(8\), which leads to potential numerical instabilities because of the correlations between successive conformations in the simulations. In a recent extension to the original WL, the statistical-temperature molecular dynamics (STMD) method \(9\), the intensive variable \(T(U) = \beta^{-1}(U)\), instead of the extensive variable \(S(U)\), is updated at each simulation step. This method is faster than the original WL, since we do not need to fill the \(N\)-order height of the curve \(S(U)\) due to the replacement. However, STMD still needs to divide the extensive variable \(U\) into order-\(N\) homogeneous bins with a prescribed bin size and it also does not sat-
Gaussian functions, like (WLL) methods cannot work unless the exact order for each phase in the system, where the Wang-Landau can visit sufficiently.

In this letter, we present a more efficient algorithm, fast adaptive flat-histogram ensemble (FAFE), to accurately estimate the DOS and enhance sampling in large general systems. FAFE calculates the ensemble means and fluctuations of $U$ using a series of generalized ensembles, then adaptively forms the points $(\beta_i, U_i)$ on the curve $\beta_i(U)$. These $U_i$ points non-uniformly distributed in $U$ space and the intervals are dependent on both $U$ and $N$ to follow the real shape of the curve $\beta_i(U)$: we automatically form many $U_i$ points in regions where $\beta_i(U)$ varies quickly, and few $U_i$ points in the regions where $\beta_i(U)$ varies slowly. A compression transformation of $U$ is also applied to reduce the visits to the $U$ regions where $\beta_i(U)$ has been well estimated. Therefore the required simulations steps of FAFE to estimate $S(U)$ are not more than $N^{1/2}$. Due to the good numerical stability of FAFE, we can visit and identify coexistent phases (including meta-stable phases) by constructing the DOS for each phase in the system, where the Wang-Landau like (WLL) methods cannot work unless the exact order parameters are known.

In a $f(U)$ ensemble, the histogram, $H_f(U) \propto e^{S(U)−f(U)}$, can be approximated as a sum of a few Gaussian functions, $H_f(U) \propto \sum_i C_i \exp\left\{-\frac{1}{2\sigma_i^2}(U−U_i)^2\right\}$, in the limit of large $N$. If $H_f(U)$ is a single-peak function, the mean of $U$ in the $f(U)$ ensemble, $\langle U \rangle_f$, is approximately equal to the maximal point of $H_f(U)$, we have,

$$\beta_s(\langle U \rangle_f) = \frac{\partial f}{\partial U}|_{\langle U \rangle_f}, \quad \frac{\partial^2 \beta_s}{\partial U^2}|_{\langle U \rangle_f} = -\sigma_f^{-2} + \frac{\partial^2 f}{\partial U^2}|_{\langle U \rangle_f},$$

where, $\sigma_f$ is the fluctuation of $U$ in the $f(U)$ ensemble. Thus, we can calculate $\langle U \rangle_f$ and $\sigma_f$ in different $f(U)$ ensembles to generate a series of points on the curve $\beta_i(U)$. The complete curve of $\beta_i(U)$ can be interpolated from these points, and $\bar{S}(U)$ can be estimated by integrating $\beta_i(U)$. This approximation, eq. (2), is sufficient in $U$ regions where simulations based on the constructed $\bar{S}(U)$ can visit sufficiently.

Different $f(U)$ ensembles, such as $f(U) = \beta U$ (canonical ensembles), and $e^{-f(U)} = (E−U)^{-1/2} \Theta(E−U)$ (microcanonical ensembles), can be used to construct $\beta_i(U)$. It is also possible to use generalized ensembles, such as

$$f(U) = \beta_0 U + \frac{1}{2}a(U−U_0)^2,$$

(3)

to generate points on $\beta_i(U)$. Because the derivative $\frac{\partial \beta_s}{\partial U}$ is positive, phase-coexistence regions are unstable in canonical ensembles. But in a generalized ensemble, e.g., $a > \frac{\partial \beta_s}{\partial U}$, they may be stabilized. Thus, once we choose suitable parameters for $(\beta_0, U_0, a)$, we can even generate the $\beta_i(U)$ data points in the phase-coexistence regions.

The implementation of FAPE is rather straightforward. Without losing any generality, we choose $U$ as the total potential energy, and define the statistical temperature $T_s(U) = \beta_s^{-1}(U)$ to illustrate it: (i) Choose an initial $T_0$ and set $T_s(U) = T_0$, then $\bar{S}(U) = U/T_0$; (ii) Simulate a segment of trajectory in the current $\bar{S}(U)$ ensemble and generate an equilibrium sample; (iii) Choose a few $f(U)$ functions (e.g. $f(U) = U/T_i$ for a few different $T_i$) and calculate the ensemble means $U_i$ and fluctuations $\sigma_i$ from the generated sample at (ii) based on the eq. (3). We choose $T_i$ from the current $T_s(U)$ function: the corresponding $U_i$, which satisfies the equation $T_i = T_s(U_i)$, must be located in the current visited regions. In addition, $|U_i − U_{i−1}| \sim \sigma_{i−1}$, where $\sigma_{i−1}$ is the fluctuation of $U$ at the temperature $T_{i−1}$. Moreover, we only need to apply the mean and fluctuation at $T_i$ when the effective size of the weighted sample, $B$, is sufficiently large. Note that we can calculate the means and fluctuations at many $T_i$ from the same sample without additional simulations. The CPU time required for calculation of the weighted mean is almost negligible compared to that for generating samples; thus we can generate more data points without much additional computational cost; (iv) From the discrete points $(\bar{U}_i, T_i)$ (and the derivative of $T_s(U)$, $T'_i = T_i^2/\sigma_i^2$), interpolate $T_s(U)$ eq. (14) and extrapolate it linearly beyond the last $U_i$. Integrate $\bar{S}(U)$ from the new $T_s(U)$; (v) Repeat (ii), (iii) and (iv) until the $T_s(U)$ and $\bar{S}(U)$ curves are complete. In Step (iii), we can check for the possible multi-peak characteristics of the weighted sample by using the generalized ensembles described by eq. (3). For example, we first choose $a = 0$ (i.e. canonical ensembles) to calculate $(T_i, \bar{U}_i)$, then set $\beta_0 = T_i^{-1}$, $U_0 = \bar{U}_i$ and use a positive $a$ to calculate the means in the new $f(U)$ ensemble, comparing the results to determine if larger $a$ should be used to depress the possible multi-peak effects. In practice, unless simulations are unexpectedly found to be trapped, this checking step is not necessary. The points $(\beta_i, U_i)$ can be calculated by using many samples generated in different segments of simulations in (ii) rather than only using a single segment of simulations, $U \leftarrow \sum U_i B_i$, where $U^j$ and $B^j$ are the the weighted mean of $U$ and the corresponding $B$ factor from the $j^{th}$ sample, respectively, and the statistical error of $\bar{U}$, which is measured by the total $B = \sum B^j$ factor, can be depressed.
Here the region is visited. The visit frequency in the large temperature region (solid phase) visit the high-$U$ region (liquid phase) visit the low-$U$ region. This compression transformation significantly decreases the computational cost of our algorithm for very large systems.

To illustrate FAFE, we apply it to truncated-and-shifted Lennard-Jones (LJ) fluids ($r_c = 2.5\sigma$) at reduced density $\rho = 0.8$. We choose $U$ as the total potential energy, and aim to calculate DOS between $T_1 = 0.5$ and $T_h = 100$, which corresponds to a very large $U$ range in large systems. The ensemble mean of $U$ is sufficiently exact if the error in $\bar{U}$ is smaller than 5 percent of the fluctuation of $U$ (i.e., $B > 400$). We apply molecular dynamics (MD) simulations with a Langevin thermostat and save a conformation every 10 MD steps. $T_s(U)$ is calculated and updated every $10^5$ steps, when the parameters of the compressive transformation are updated. In our simulations, the distance between neighboring $U_i$ points is far larger than that in WLL [9] in high-$T$ regions and large-$N$ systems, but automatically decreases in low $T$ regions and small $N$ systems. Figure (1) shows that $T_s(U/N)$, $c_{sv} = (dT_s/d(U/N))^{-1}$ and $s(u) = (S(U)/N)$ in systems with several different $N$ values are quickly generated in the large temperature regions. The curves for different $N$ all collapse into single curves as expected. Figure (2) shows the efficiency of FAFE, the number of MD steps $t \propto N^\gamma$ with $\gamma \approx 0.5$.

As shown in Fig. (3), in the lower temperature region, $T_s(U)$ has multiple branches that correspond to the DOS of some separated parts in conformational space. For a segment of simulated trajectory, we may calculate the corresponding $T_s(U)/T$ points, and learn which branch of $T_s(U)$ the points are located on (i.e., which macroscopic state the segment of trajectory resides in). The trajectories generated under the high-$U$ branch of $T_s(U)$ (liquid phase) have the expected flat histograms in the region $U > U_1$, but quickly transition into the low-$U$ region (solid phase). Similarly, simulations under the low-$U$ branch of $T_s(U)$ (solid phase) visit the $U < U_2$ region with equal probabilities, but transition into the liquid phase. The overlap of the two branches indicates...
that the potential energy of the liquid phase can be lower than that of the solid phase, and $U$ is definitely not the proper order parameter of the liquid/solid phase transition. It is very hard to visit (and more difficult to identify) the low-energy liquid and high-energy solid conformations in current existing simulation techniques. Thus, FAFE, which is able to visit more parts of the configuration space and identify them by forming their own DOS without requiring the prior knowledge of precise order parameters, might provide new promise in understanding phase transition processes.

It is possible to calculate the total $S(U)$ and $T_s(U)$ from these obtained $T_s(U)$ branches [10]. However, even in the exact $S(U)$ ensemble, it is still not guaranteed that simulations can freely transition between the states. In this case, the visited probability of conformation inside each iso-$U$ is constant, $P(r) \propto e^{-S(U)}$, then any small-fraction part in an iso-$U$ subspace is still rarely visited although the total visited probability of the iso-$U$ subspace is not small. If a small-fraction part corresponds to the passage between two states, simulations in the $S(U)$ ensemble still do not easily pass the passage. The $S(U)$ ensemble only provides chances to search transition passages in more iso-$U$ subspaces. This is a general limitation in all the low-dimension flat-histogram ensemble methods. The algorithms with good stabilities (e.g., simulations satisfying detailed balance) are helpful in analyzing and (partly) overcoming the limitation. For example, FAFE generates the $T_s(U)$ and $S(U)$ (with arbitrary constant) in each visited region instead of the expected total $T_s(U)$ and $S(U)$ of the whole conformational space. Thus, even when the equilibration among the regions is difficult, it is still possible to provide many useful results. From the branches of $S(U)$ or $T_s(U)$, it is possible to construct the $S(U)$ for whole the conformational space [10]. In addition, we can apply the generalized ensembles described by eq. [3] to find more (possibly metastable) states. For example, besides the stable liquid and solid phases in the LJ system, we detect several metastable solid states with different crystalline structures as well as the unstable multi-phase coexistence, which are shown in Fig.(4). These complex solid phases arise from the fact that the system is fixed at $\rho = 0.8$ so the interparticle distance in the face-center cubic structure does not match the equilibrium distance of LJ potential, $r_0 = 2^{1/6}$.

The flat-histogram ensembles based on multiple collective variables are expected to be better for enhanced sampling than single variable [10, 12]. The required computational cost will be $O(N^d)$, where $d$ is the number of the collective variables. FAFE ($\gamma = 1/2$), which is much faster than WLL ($\gamma = 1.5 \sim 3$), is efficient in sampling large complex systems, such as glasses and proteins. It is useful to construct the canonical DOS $\Omega(U; T) = \int e^{-V(r)/T} \delta(U - V(r)) dr$ instead of the normal DOS in FAFE in order to keep the connectivity of molecules and more focus on the desired temperature $T$, if the total potential energy is not chosen as $U$. It is also possible to combine FAFE with the replica exchange method (REM): we generate the branches of $T_s(U)$ and use them in different replicas and exchange conformations among these replicas; thus we can use $O(N^0)$ replicas in large systems instead of the $N^{1/2}$—order replicas required in conventional REM, which is the main limitation of applying REM.

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[13] In the weighted sample, the fluctuation of $Q$ is calculated as,
$$\sigma^2 = \frac{\sum w_k (\sum w_k Q^2) - (\sum w_k Q)^2}{\sum w_k}$$

[14] High-order interpolation is also possible by using $T'_i$.
[15] In practice, we linearly extrapolate $T_s(U)$ in a finite range from the previous $U_i$ point, and set $T_s(U)$ as a constant out of the range.
[16] The total entropy $S(U) = \ln \sum_i \exp[S^i(U)]$. $S^i(U) = S^i_0 + \int_{U_i}^U T^i_s(V) dV$ and $T^i_s(U)$ are the $i^{th}$ branch of the entropy and temperature, respectively.