# A simple method for automated equilibration detection in molecular simulations

John D. Chodera<sup>1, \*</sup>

<sup>1</sup>Computational Biology Program, Sloan Kettering Institute, Memorial Sloan Kettering Cancer Center, New York, NY 10065 (Dated: March 7, 2016)

Molecular simulations intended to compute equilibrium properties are often initiated from configurations that are highly atypical of equilibrium samples, a practice which can generate a distinct initial transient in mechanical observables computed from the simulation trajectory. Traditional practice in simulation data analysis recommends this initial portion be discarded to *equilibration*, but no simple, general, and automated procedure for this process exists. Here, we suggest a conceptually simple automated procedure that does not make strict assumptions about the distribution of the observable of interest, in which the equilibration time is chosen to maximize the number of effectively uncorrelated samples in the production timespan used to compute equilibrium averages. We present a simple Python reference implementation of this procedure, and demonstrate its utility on typical molecular simulation data.

Keywords: molecular dynamics (MD); Metropolis-Hastings; Monte Carlo (MC); Markov chain Monte Carlo (MCMC); equilibration; burn-in; timeseries analysis; statistical inefficiency; integrated autocorrelation time

## INTRODUCTION

Molecular simulations use Markov chain Monte Carlo (MCMC) techniques [1] to sample configurations x from an equilibrium distribution  $\pi(x)$ , either exactly (using Monte Carlo methods such as Metropolis-Hastings) or approximately (using molecular dynamics integrators without Metropolization) [2].

Due to the sensitivity of the equilibrium probability density  $\pi(x)$  to small perturbations in configuration x and the difficulty of producing sufficiently good guesses of typical equilibrium configurations  $x \sim \pi(x)$ , these molecular simulations are often started from highly atypical initial conditions. For example, simulations of biopolymers might be initiated from a fully extended conformation unrepresentative of behavior in solution, or a geometry derived from a fit to diffraction data collected from a cryocooled crystal; solvated systems may be prepared by periodically replicating small solvent box equilibrated under different conditions, yielding atypical densities and solvent structure; liquid mixtures or lipid bilayers may be constructed by using methods that fulfill spatial constraints (e.g. PackMol [3]) but create locally aytpical geometries, requiring long simulation times to relax to typical configurations.

As a result, traditional practice in molecular simulation has recommended some initial portion of the trajectory be discarded to *equilibration* (also called *burn-in*<sup>1</sup> in the MCMC literature [4]). While the process of discarding initial samples is strictly unnecessary for the time-average of quantities of interest to eventually converge to the desired expectations [5], this nevertheless often allows the practitioner to avoid what may be impractically long run times to eliminate the bias in computed properties in finite-length simulations

38 induced by atypical initial starting conditions. It is worth
39 noting that a similar procedure is not a practice universally
40 recommended by statisticians when sampling from poste41 rior distributions in statistical inference [4]; the differences
42 in complexity of probability densities typically encountered
43 in statistics and molecular simulation may explain the dif44 ference in historical practice.

As a motivating example, consider the computation of the 46 average density of liquid argon under a given set of reduced 47 temperature and pressure conditions shown in Figure 1. To 48 initiate the simulation, an initial dense liquid geometry at <sub>49</sub> reduced density  $\rho^* \equiv \rho \sigma^3 = 0.960$  was prepared and sub-50 jected to local energy minimization. The upper panel of Fig-51 ure 1 depicts the average relaxation behavior of simulations 52 initiated from the same configuration with different random 53 initial velocities and integrator random number seeds (see 54 Simulation Details). The average of 500 realizations of this 55 process shows a characteristic relaxation away from the ini-56 tial density toward the equilibrium density (Figure 1, upper 57 panel, black line). As a result, the expectation of the run-58 ning average of the density significantly deviates from the 59 true expectation (Figure 1, lower panel, dashed line). This 60 effect leads to significantly biased estimates of the expec-61 tation unless simulations are sufficiently long to eliminate 52 starting point dependent bias, which takes a surprisingly  $_{63}$  long  $\sim$ 2000au in this example. Note that this bias is present even in the average of many realizations because the same 65 atypical starting condition is used for every realization of 66 this simulation process.

To develop an automatic approach to eliminating this bias, we take motivation from the concept of *reverse cumulative averaging* from Yang et al. [6], in which the trajectory statistics over the production region of the trajectory are examined for different choices of the end of the discarded equilibration region to determine the optimal production region to use for computing expectations and other statistical properties. We begin by first formalizing our objectives mathematically.

<sup>\*</sup> Corresponding author; john.chodera@choderalab.org

<sup>&</sup>lt;sup>1</sup> The term *burn-in* comes from the field of electronics, in which a short "burn-in" period is used to ensure that a device is free of faulty components—which often fail quickly—and is operating normally [4].

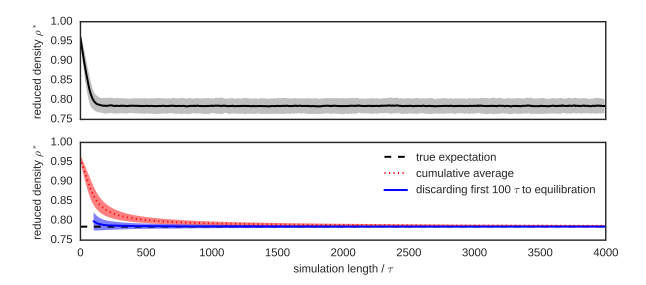


FIG. 1. Illustration of the motivation for discarding data to equilibration. To illustrate the bias in expectations induced by relaxation away from initial conditions, 500 replicates of a simulation of liquid argon were initiated from the same energy-minimized initial configuration constructed with initial reduced density  $\rho^* \equiv \rho \sigma^3 = 0.960$  but different random number seeds for stochastic integration. **Top:** The average of the reduced density (black line) over the replicates relaxes to the region of typical equilibrium densities over the first  $\sim 100~\tau$  of simulation time, where  $\tau$  is a natural time unit (see *Simulation Details*). **Bottom:** If the average density is estimated by a cumulative average from the beginning of the simulation (red dotted line), the estimate will be heavily biased by the atypical starting density even beyond  $1000~\tau$ . Discarding even a small amount of initial data—in this case 500 initial samples—results in a cumulative average estimate that converges to the true average (black dashed line) much more rapidly. Shaded regions denote 95% confidence intervals.

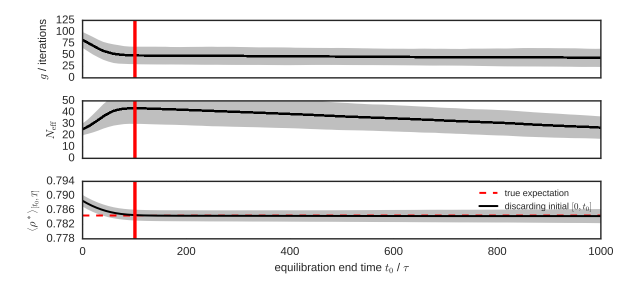


FIG. 2. Statistical inefficiency, number of uncorrelated samples, and bias for different equilibration times. Trajectories of length  $T=2000~\tau$  for the argon system described in Figure 1 were analyzed as a function of equilibration time choice  $t_0$ . Averages over all 500 replicate simulations (all starting from the same initial conditions) are shown as dark lines, with shaded lines showing standard deviation of estimates among replicates. **Top:** The statistical inefficiency g as a function of equilibration time choice  $t_0$  is initially very large, but diminishes rapidly after the system has relaxed to equilibrium. **Middle:** The number of effectively uncorrelated samples  $N_{\rm eff} = (T-t_0+1)/g$  shows a maximum at  $t_0 \sim 100~\tau$  (red vertical lines), suggesting the system has equilibrated by this time. **Bottom:** The cumulative average density  $\langle \rho^* \rangle$  computed over the span  $[t_0, T]$  shows that the bias (deviation from the true estimate, shown as red dashed lines) is minimized for choices of  $t_0 \geq 100~\tau$ . The standard deviation among replicates (shaded region) grows with  $t_0$  because fewer data are included in the estimate. The choice of optimal  $t_0$  that maximizes  $N_{\rm eff}$  (red vertical line) strikes a good balance between bias and variance. The true estimate (red dashed lines) is computed from averaging over the range  $[5~000,10~000]~\tau$  over all 500 replicates.

#### STATEMENT OF THE PROBLEM

Consider T successively sampled configurations  $x_t$  from a molecular simulation, with  $t=1,\ldots,T$ , initiated from  $x_0$ . By We presume we are interested in computing the expectation

$$\langle A \rangle \equiv \int dx \, A(x) \, \pi(x)$$
 (1) 103

 $_{\text{80}}$  of a mechanical property of interest A(x). For convenience,  $_{\text{81}}$  we will refer to the timeseries  $a_t\equiv A(x_t),$  with  $t\in[1,T].$  The estimator  $\hat{A}\approx\langle A\rangle$  constructed from the entire dataset  $_{\text{83}}$  is given by

$$\hat{A}_{[1,T]} \equiv \frac{1}{T} \sum_{t=1}^{T} a_t.$$
 (2)

 $^{\rm 84}$  While  $\lim_{T\to\infty}\hat{A}_{[1,T]}=\langle A\rangle$  for an infinitely long simulass tion², the bias in  $\hat{A}_{[1,T]}$  may be significant in a simulation of finite length T.

By discarding samples  $t < t_0$  to equilibration, we hope to exclude the initial transient from our sample average, and provide a less biased estimate of  $\langle A \rangle$ ,

$$\hat{A}_{[t_0,T]} \equiv \frac{1}{T - t_0 + 1} \sum_{t=t_0}^{T} a_t. \tag{3}$$

We can quantify the overall error in an estimator  $A_{[t_0,T]}$  in a sample average of trajectories initiated from  $x_0$  that excludes samples where  $t < t_0$  by the expected error  $\delta^2 \hat{A}_{[t_0,T]}$ ,

$$\delta^2 \hat{A}_{[t_0,T]} \equiv E_{x_0} \left[ \left( \hat{A}_{[t_0,T]} - \langle A \rangle \right)^2 \right] \tag{4}$$

 $_{^{94}}$  where  $E_{x_0}[\cdot]$  denotes the expectation over independent re-  $_{^{95}}$  alizations of the specific simulation process initiated from  $_{^{96}}$  configuration  $x_0$ , but with different velocities and random  $_{^{97}}$  number seeds.

 $^{98}$   $\,$  We can rewrite the expected error  $\delta^2 \hat{A}$  by separating it  $^{99}$  into two components:

$$\delta^{2} \hat{A}_{[t_{0},T]} = E_{x_{0}} \left[ \left( \hat{A}_{[t_{0},T]} - E_{x_{0}} [\hat{A}_{[t_{0},T]}] \right)^{2} \right] + \left( E_{x_{0}} [\hat{A}_{[t_{0},T]}] - \langle A \rangle \right)^{2}$$
(5)

The first term denotes the variance in the estimator  $\hat{A}$ ,

$$\operatorname{var}_{x_0}(\hat{A}_{[t_0,T]}) \equiv E_{x_0} \left[ \left( \hat{A}_{[t_0,T]} - E_{x_0}[\hat{A}_{[t_0,T]}] \right)^2 \right]$$
 (6)

while the second term denotes the contribution from the squared bias,

$$bias_{x_0}^2(\hat{A}_{[t_0,T]}) \equiv \left(E_{x_0}[\hat{A}_{[t_0,T]}] - \langle A \rangle\right)^2 \tag{7}$$

#### **BIAS-VARIANCE TRADEOFF**

With increasing equilibration time  $t_0$ , bias is reduced, but T].

with evariance—the contribution to error due to random variation time  $t_0$ , bias is reduced, but the variance—the contribution to error due to random variation time from having a finite number of uncorrelated samples—will increase because less data is included in the estimate.

This can be seen in the bottom panel of Figure 2, where the shaded region (95% confidence interval of the mean) increases in width with increasing equilibration time  $t_0$ .

To examine the tradeoff between bias and variance explicitly, Figure 3 plots the bias and variance (here, shown as the standard deviation over replicates—the square root of the variance—which is an indication of the true standard error of a single simulation) contributions against each other as a function of  $t_0$  (denoted by color) as computed from statistics over all 500 replicates. At  $t_0=0$ , the bias is large but variance is minimized. With increasing  $t_0$ , bias is eventually eliminated but then variance rapidly grows as fewer uncorrelated samples are included in the estimate. There is a clear optimal choice at  $t_0\sim 100\,\tau$  that minimizes variance while also effectively eliminating bias (where  $\tau$  is a natural time unit—see  $Simulation\ Details$ ).

# **SELECTING THE EQUILIBRATION TIME**

ls there a simple approach to choosing an optimal equilibration time  $t_0$  that provides a significantly improved estimate  $\hat{A}_{[t_0,T]}$ , even when we do not have access to multiple realizations? At worst, we hope that such a procedure would at least give some improvement over the naive estimate, such that  $\delta^2\hat{A}_{[t_0,T]}<\delta^2\hat{A}_{[0,T]}$ ; at best, we hope that we can achieve a reasonable bias-variance tradeoff close to the optimal point identified in Figure 3 that minimizes bias without greatly increasing variance. We remark that, for cases in which the simulation is not long enough to reach equilibrium, no choice of  $t_0$  will eliminate bias completely; the best we can hope for is to minimize this bias.

While automated methods for selecting the equilibration time  $t_0$  have been proposed, these approaches have short-comings that have greatly limited their use. The reverse tumulative averaging (RCA) method proposed by Yang et al. [6], for example, uses a statistical test for normality to determine the point before which which the observable timeseries deviates from normality when examining the timeseries in reverse. While this concept may be reasonable for experimental data, where measurements often represent the sum of many random variables such that the central limit theorem's guarantee of asymptotic normality ensures the distribution of the observable will be approximately normal, there is no such guarantee that instantaneous mea-

 $<sup>^2</sup>$  We note that this equality only holds for simulation schemes that sample from the true equilibrium density  $\pi(x)$ , such as Metropolis-Hastings Monte Carlo or Metropolized dynamical integration schemes such as hybrid Monte Carlo (HMC). Molecular dynamics simulations utilizing finite timestep integration without Metropolization will produce averages that may deviate from the true expectation  $\langle A \rangle$  [2].

mally distributed. In fact, many properties will be decidedly  $_{183}$  of the sampling interval  $\tau$ ) are given by non-normal. For a biomolecule such as a protein, for example, the radius of gyration, end-to-end distance, and torsion angles sampled during a simulation will all be highly nonnormal. Instead, we require a method that makes no assumptions about the nature of the distribution of the property under study.

#### **AUTOCORRELATION ANALYSIS**

158

The set of successively sampled configurations  $\{x_t\}$  and their corresponding observables  $\{a_t\}$  compose a correlated timeseries of observations. To estimate the statistical error or uncertainty in a stationary timeseries free of bias, we must be able to quantify the effective number of uncorrelated samples present in the dataset. This is usually accomplished through computation of the statistical ineffiiency g, which quantifies the number of correlated timeeries samples needed to produce a single effectively uncorrelated sample of the observable of interest. While these concepts are well-established for the analysis of both Monte Carlo and molecular dynamics simulations [7-10], we review them here for the sake of clarity.

For a given equilibration time choice  $t_0$ , the statistical uncertainty in our estimator  $\hat{A}_{[t_0,T]}$  can be written as,

$$\delta^{2} \hat{A}_{[t_{0},T]} \equiv E_{x_{0}} \left[ \left( \hat{A}_{[t_{0},T]} - \langle \hat{A} \rangle \right)^{2} \right]$$

$$= E_{x_{0}} \left[ \hat{A}_{[t_{0},T]}^{2} \right] - E_{x_{0}} \left[ \hat{A}_{[t_{0},T]} \right]^{2}$$

$$= \frac{1}{T_{t_{0}}^{2}} \sum_{t,t'=t_{0}}^{T} \left\{ E_{x_{0}} \left[ a_{t} a_{t'} \right] - E_{x_{0}} \left[ a_{t} \right] E_{x_{0}} \left[ a_{t'} \right] \right\}$$

$$= \frac{1}{T_{t_{0}}^{2}} \sum_{t=t_{0}}^{T} \left\{ E_{x_{0}} \left[ a_{t}^{2} \right] - E_{x_{0}} \left[ a_{t} \right]^{2} \right\}$$

$$+ \frac{1}{T_{t_{0}}^{2}} \sum_{t=t_{0}}^{T} \left\{ E_{x_{0}} \left[ a_{t} a_{t'} \right] - E_{x_{0}} \left[ a_{t} \right] E_{x_{0}} \left[ a_{t'} \right] \right\},$$

$$(8)$$

where  $T_{t_0}\equiv T-t_0+1$ , the number of correlated samples in the timeseries  $\{a_t\}_{t_0}^T$ . In the last step, we have split the double-sum into two separate sums—a term capturing the variance in the observations  $a_t$ , and a remaining term capturing the correlation between observations.

If  $t_0$  is sufficiently large for the initial bias to be eliminated, the remaining timeseries  $\{a_t\}_{t_0}^T$  will obey the properties of both stationarity and time-reversibility, allowing us to write,

$$\delta^{2} \hat{A}_{[t_{0},T]}^{\text{equil}} = \frac{1}{T_{t_{0}}} \left[ \langle a_{t}^{2} \rangle - \langle a_{t} \rangle^{2} \right]$$

$$+ \frac{2}{T_{t_{0}}} \sum_{n=1}^{T-t_{0}} \left( \frac{T_{t_{0}} - n}{T_{t_{0}}} \right) \left[ \langle a_{t} a_{t+n} \rangle - \langle a_{t} \rangle \langle a_{t+n} \rangle \right]$$

$$\equiv \frac{\sigma_{t_{0}}^{2}}{T_{t_{0}} / a_{t_{0}}}, \tag{9}$$

<sub>150</sub> surements of a simulation property of interest will be nor-  $_{182}$  where the variance  $\sigma^2$  and statistical inefficiency g (in units

$$\sigma^2 \equiv \langle a_t^2 \rangle - \langle a_t \rangle^2, \tag{10}$$

$$g \equiv 1 + 2\tau_{\rm eq},\tag{11}$$

integrated autocorrelation time  $au_{
m ac}$  given by

$$\tau_{\rm eq} \equiv \sum_{t=1}^{T-1} \left( 1 - \frac{t}{T} \right) C_t, \tag{12}$$

(13)

with the discrete-time normalized fluctuation autocorrela-186 tion function  $C_t$  defined as

$$C_t \equiv \frac{\langle a_n a_{n+t} \rangle - \langle a_n \rangle^2}{\langle a_n^2 \rangle - \langle a_n \rangle^2}.$$
 (14)

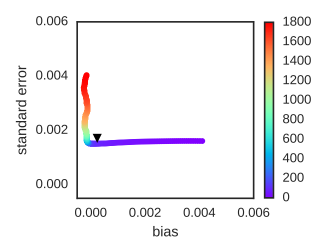
In practice, it is difficult to estimate  $C_t$  for  $t \sim T$ , due to  $_{188}$  growth in the statistical error, so common estimators of g $_{189}$  make use of several additional properties of  $C_t$  to provide useful estimates (see Practical Computation of Statistical In-191 efficiencies).

The  $t_0$  subscript for the variance  $\sigma^2$ , the integrated autocorrelation time  $au_{\mathrm{ac}}$ , and the statistical inefficiency g mean that these quantities are only estimated over the production portion of the timeseries,  $\{a_t\}_{t=t_0}^T$ . Since we assumed that 196 the bias was eliminated by judicious choice of the equilibra- $_{\scriptscriptstyle{197}}$  tion time  $t_0$  , this estimate of the statistical error will be poor  $_{198}$  for choices of  $t_0$  that are too small.

# THE ESSENTIAL IDEA

Suppose we choose some arbitrary time  $t_0$  and discard all samples  $t \in [0, t_0)$  to equilibration, keeping  $[t_0, T]$  as the dataset to analyze. How much data remains? We can determine this by computing the statistical inefficiency  $g_{t_0}$  for the interval  $[t_0, T]$ , and computing the effective number of uncorrelated samples  $N_{\rm eff}(t_0) \equiv (T-t_0+1)/g_{t_0}$ . If we start at  $t_0 \equiv T$  and move  $t_0$  to earlier and earlier points in time, we expect that the effective number of uncorrelated samples  $N_{\rm eff}(t_0)$  will continue to grow until we start to include the highly atypical initial data. At that point, the integrated autocorrelation time au (and hence the statistical inefficiency g) will greatly increase (a phenomenon observed 212 earlier, e.g. Figure 2 of [6]). As a result, the effective number of samples  $N_{\mathrm{eff}}$  will start to plummet.

Figure 2 demonstrates this behavior for the liquid argon system described above, using averages of the statistical inefficiency  $g_{t_0}$  and  $N_{\mathrm{eff}}(t_0)$  computed over 500 independent replicate trajectories. At short  $t_0$ , the average statistical inefficiency g (Figure 2, top panel) is large due to the contribution from slow relaxation from atypical initial conditions, while at long  $t_0$  the statistical inefficiency estimate is much shorter and nearly constant of a large span of time 222 origins. As a result, the average effective number of uncor- $_{
m 223}$  related samples  $N_{
m eff}$  (Figure 2, middle panel) has a peak at  $_{
m 224}$   $t_0\sim 100\, au$  (Figure 2, vertical red lines). The effect on bias in



Bias-variance tradeoff for fixed equilibration time versus automatic equilibration time selection. Trajectories of length  $T=2000\tau$  for the argon system described in Figure 1 were analyzed as a function of equilibration time choice  $t_0$ , with colors denoting the value of  $t_0$  (in units of  $\tau$ ) corresponding to each plotted point. Using 500 replicate simulations, the average bias (average deviation from true expectation) and standard deviation (random variation from replicate to replicate) were computed as a function of a prespecified fixed equilibration time  $t_0$ , with colors running from violet  $(0 \tau)$  to red  $(1800 \tau)$ . As is readily discerned, the bias for small  $t_0$  is initially large, but minimized for larger  $t_0$ . By contrast, the standard error (a measure of variance, estimated here by standard deviation among replicates) grows as  $t_0$  grows above a certain critical time (here,  $\sim 100 \, \tau$ ). If the  $t_0$  that maximizes  $N_{\rm eff}$ is instead chosen individually for each trajectory based on that trajectory's estimates of statistical inefficiency  $g_{[t_0,T]}$  , the resulting  $_{^{264}}$ bias-variance tradeoff (black triangle) does an excellent job minimizing bias and variance simultaneously, comparable to what is possible for a choice of equilibration time  $t_0$  based on knowledge of the true bias and variance among many replicate estimates.

the estimated average reduced density  $\langle \rho^* \rangle$  (Figure 2, bottom panel) is striking—the bias is essentially eliminated for the choice of equilibration time  $t_0$  that maximizes the number of uncorrelated samples  $N_{\rm eff}$ .

This suggests an alluringly simple algorithm for identify- $_{
m 230}$  ing the optimal equilibration time—pick the  $t_0$  which maxi- $_{
m 231}$  mizes the number of uncorrelated samples  $N_{
m eff}$  in the time-232 series  $\{a_t\}_{t_0}^T$  for the quantity of interest A(x):

$$t_0^{\text{opt}} = \underset{t_0}{\operatorname{argmax}} N_{\text{eff}}(t_0)$$

$$= \underset{t_0}{\operatorname{argmax}} \frac{T - t_0 + 1}{g_{t_0}}$$
(15)

234 selecting the equilibration time  $t_0$  using Eq 15 work for cases 285 ferent choice of observable A(x) for the same liquid argon

where we do not know the statistical inefficiency g as a function of the equilibration time  $t_0$  precisely? When all that is available is a single simulation, our best estimate of  $g_{t_0}$  is derived from that simulation alone over the span  $[t_0, T]$  will this affect the quality of our estimate of equilibration 240 time? Empirically, this does not appear to be the case the black triangle in Figure 3 shows the bias and variance 242 contributions to the error for estimates computed over the  $_{243}$  500 replicates where  $t_0$  is individually determined from each  $_{\mbox{\tiny 244}}$  simulation using this simple scheme based on selecting  $t_0$ to maximize  $N_{
m eff}$  for each individual realization. Despite not <sup>246</sup> having knowledge about multiple realizations, this strategy 247 effectively achieves a near-optimal balance between minimizing bias without increasing variance.

Overall RMS error. How well does this strategy perform  $_{ extstyle 250}$  in terms of decreasing the  $\mathit{overall}$  error  $\delta \hat{A}_{[t_0,T]}$  compared to  $\delta \hat{A}_{[0,T]}$ ? Figure 4 compares the expected standard error (de-252 noted  $\delta \hat{A}$ ) as a function of a fixed initial equilibration time  $t_0$ 253 (black line with shaded region denoting 95% confidence in- $_{ ext{254}}$  terval) with the strategy of selecting  $t_0$  to maximize  $N_{
m eff}$  for each realization (red line with shaded region denoting 95% confidence interval). While the minimum error for the fixed $t_0$  strategy (0.00152 $\pm$ 0.00005) is achieved at  $\sim 100 \, \tau$ —a fact that could only be determined from knowledge of multiple realizations—the simple strategy of selecting  $t_0$  using Eq. 15 260 achieves a minimum error of 0.00173±0.00005, only 11% worse (compared to errors of 0.00441 $\pm$ 0.00007, or 290% worse, should no data have been discarded).

# **DISCUSSION**

The scheme described here—in which the equilibration  $time t_0$  is computed using Eq. 15 as the choice that maximizes the number of uncorrelated samples in the produc $tion region [t_0, T]$ —is both conceptually and computation-268 ally straightforward. It provides an approach to determin-269 ing the optimal amount of initial data to discard to equilibration in order to minimize variance while also minimizing initial bias, and does this without employing statistical tests that require generally unsatisfiable assumptions of normal-273 ity of the observable of interest. All that is needed is to save  $_{ extstyle 274}$  the timeseries  $\{a_t\}_1^T$  of the observable A(x) of interest there is no need to store full configurations  $x_t$ —and postprocess this dataset with a simple analysis procedure, for which we have provided a convenient Python reference implementation (see Simulation Details). As we have seen, this scheme empirically appears to select a practical compromise between bias and variance even when the statistical inefficiency q is estimated directly from the trajectory using 282 Eq. 11.

To show that this approach is indeed general, we re-Bias-variance tradeoff. How will the simple strategy of 284 peated the analysis illustrated above in Figs. 1-4 for a dif-

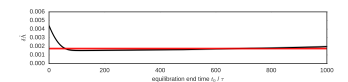


FIG. 4. RMS error for fixed equilibration time versus automatic equilibration time selection. Trajectories of length  $T=2000\tau$  for the argon system described in Figure 1 were analyzed as a function of fixed equilibration time choice  $t_0$ . Using 500 replicate simulations, the root-mean-squared (RMS) error (Eq. 4) was computed (black line) along with 95% confidence interval (gray shading). The RMS error is minimized for fixed equilibration time choices in the range 100–200  $\tau$ . If the  $t_0$  that maximizes  $N_{\rm eff}$  is instead chosen individually for each trajectory based on that trajectory's estimated statistical inefficiency  $g_{[t_0,T]}$  using Eq. 15, the resulting RMS error (red line, 95% confidence interval shown as red shading) is quite close to the minimum RMS error achieved from any particular fixed choice of equilibration time  $t_0$ , suggesting that this simple automated approach to selecting  $t_0$  achieves close to optimal performance.

tion time  $t_0$  from Eq. 15 does just as well.

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estimated autocorrelation time.

#### SIMULATION DETAILS

All molecular dynamics simulations described here were performed with OpenMM 6.3 [12] (available at openmm.org) using the Python API. All scripts used to retrieve the software versions used here, run the simulations, analyze data, and generate plots—along with the simulation data itself and scripts for generating figures—are available on GitHub<sup>4</sup>.

To model liquid argon, the LennardJonesFluid model 313 system in the openmmtools package<sup>5</sup> was used with pa-<sub>314</sub> rameters appropriate for liquid argon ( $\sigma=$  3.4 Å,  $\epsilon=$ 

system—in this case, the reduced potential energy  $^3u^*(x) \equiv _{315}$  0.238 kcal/mol). All results are reported in reduced (dimen- $\beta U(x)$ . The results of this analysis are collected in Fig. 5. As 316 sionless) units. Initial dense liquid geometries were gencan readily be seen, this reduced potential behaves in es- 317 erated via a Sobol' subrandom sequence [? ], as genersentially the same way the reduced density does, and the 318 ated by the subrandom\_particle\_positions method in simple scheme for automated determination of equilibra- 319 openmmtools. A cubic switching function was employed, with the potential gently switched to zero over  $r \in [\sigma, 3\sigma]$ , A word of caution is necessary. One can certainly envision 321 and a long-range isotropic dispersion correction accounting pathological scenarios where this algorithm for selecting an 322 for this switching behavior used to include neglected contrioptimal equilibration time will break down. In cases where  $_{
m 323}$  butions. Simulations were performed using a periodic box the simulation is not long enough to reach equilibrium—let  $^{324}$  of N=500 atoms at reduced temperature  $T^*\equiv k_BT/\epsilon=100$ alone collect many uncorrelated samples from it—no choice  $_{325}$  0.850 and reduced pressure  $p^*\equiv p\sigma^3/\epsilon=1.266$  using a of equilibration time will bestow upon the experimenter the  $_{326}$  Langevin integrator [13] with timestep  $\Delta t = 0.01 au$  and colability to produce an unbiased estimate of the true expecta- 327 lision rate  $\nu = \tau^{-1}$ , with characteristic oscillation timescale tion. Similarly, in cases where insufficient data is available  $\tau = \sqrt{mr_0^2/72\epsilon}$  and  $\tau = 2^{1/6}\sigma$  [14]. All times are reported for the statistical inefficiency to be estimated well, this al-  $\frac{329}{2}$  in multiples of the characteristic timescale  $\tau$ . A molecugorithm is expected to perform poorly. However, in these 330 lar scaling Metropolis Monte Carlo barostat with Gaussian cases, the data itself should be suspect if the trajectory is 331 simulation volume change proposal moves attempted evnot at least an order of magnitude longer than the minimum  $^{332}$  ery au (100 timesteps), using an adaptive algorithm that adjusts the proposal width during the initial part of the simulation [12]. Densities were recorded every  $\tau$  (100 timesteps). 335 The true expectation  $\langle \rho^* \rangle$  was estimated from the sample average over all 500 realizations over [5000,10000]  $\tau$ .

> The automated equilibration detection scheme is also available in the timeseries module of the pymbar package as detectEquilibration(), and can be accessed us-340 ing the following code:

from pymbar.timeseries import detectEquilibration # determine equilibrated region [t0, g, Neff\_max] = detectEquilibration(A\_t) # discard initial samples to equilibration  $A_t = A_t[t0:]$ 

# PRACTICAL COMPUTATION OF STATISTICAL INEFFICIENCIES

The robust computation of the statistical inefficiency q 343 (defined by Eq. 11) for a finite timeseries  $a_t$ ,  $t=0,\ldots,T$ 344 deserves some comment. There are, in fact, a variety of http://github.com/choderalab/automatic-equilibration-deteggischemes for estimating q described in the literature, and their behaviors for finite datasets may differ, leading to dif-

<sup>&</sup>lt;sup>3</sup> Note that the reduced potential [11] for the isothermal-isobaric ensemble is generally defined as  $u^*(x) = \beta[u(x) + pV(x)]$  to include the pressure-volume term  $\beta pV(x)$ , but in order to demonstrate the performance of this analysis on an observable distinct from the density—which depends on V(x)—we omit the  $\beta pV(x)$  term in the present analysis.

<sup>&</sup>lt;sup>4</sup> All Python scripts necessary to reproduce this work—along with data plotted in the published version—are available at:

<sup>&</sup>lt;sup>5</sup> available at http://github.com/choderalab/openmmtools

 $_{347}$  ferent estimates of the equilibration time  $t_0$  using the algo- $_{373}$  variant of the *initial positive sequence* (IPS) method of Geyer rithm of Eq. 15.

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The main issue is that a straightforward approach to estimating the statistical inefficiency using Eqs. 12-14 in which the expectations are simply replaced with sample estimates auses the statistical error in the estimated correlation function  $C_t$  to grow with t in a manner that allows this error to quickly overwhelm the sum of Eq. 12. As a result, a number of alternative schemes—generally based on controlling the error in the estimated  $C_t$  or truncating the sum of Eq. 12 when the error grows too large—have been proposed.

For stationary, irreducible, reversible Markov chains, Geyer observed that a function  $\Gamma_k \equiv \gamma_{2k} + \gamma_{2k+1}$  of the unnormalized fluctuation autocorrelation function  $\gamma_t \equiv$  $\langle a_i a_{i+t} \rangle - \langle a_i \rangle^2$  has a number of pleasant properties (Theorem 3.1 of [15]): It is strictly positive, strictly decreasing, and strictly convex. Some or all of these properties can be exploited to define a family of estimators called initial sequence methods (see Section 3.3 of [15] and Section 1.10.2 of [4]), of which the initial convex sequence (ICS) estimator is generally agreed to be optimal, if somewhat more complex to implement.6 368

(data not shown). This method is related to a multiscale 396 course of this work.

<sub>374</sub> [16], where contributions are accumulated at increasingly longer lag times and the sum of Eq. 12 is truncated when the terms become negative. We have found this method to be both fast and to provide useful estimates of the statistical inefficiency, but it may not perform well for all problems.

## **ACKNOWLEDGMENTS**

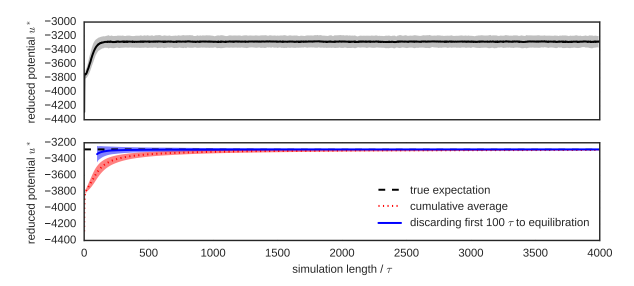
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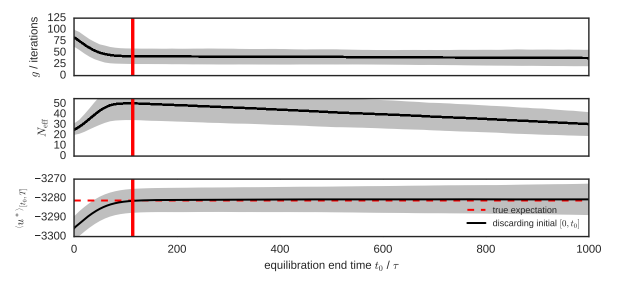
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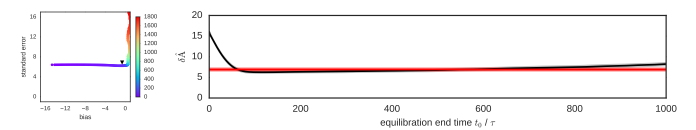
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(a) *Top*: Average reduced potential for relaxation from initial conditions (black solid line). *Bottom*: Cumulative averages of reduced potential with (blue solid line) and without (red dotted line) discarding some initial data to equilibration.



(b) Statistical inefficiency g, effective number of uncorrelated samples  $N_{\rm eff}$ , and average of reduced potential  $u^*$  omitting equilibration time  $t_0$  for analysis of reduced potential.



(c) Bias-variance tradeoff for (d) Overall RMSE error in estimate of reduced potential as a function of discarded equilibration time (black) compared reduced potential. with RMSE from automatic equilibration detection scheme (red line).

FIG. 5. Corresponding analysis for reduced potential energy of liquid argon system. The analyses of Figs. 1–4 were repeated for the reduced potential energy  $u^*(x) \equiv \beta U(x)$  of the liquid argon system. As with the analysis of reduced density, the simple automated determination of equilibration time  $t_0$  from Eq. 15 works equivalently well for the reduced potential. Shaded regions denote 95% confidence interval.