[DRAFT]: Best Practices for Thermodynamic Property Prediction from Molecular Simulations

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This document describes a collected set of best practices for computing various physical properties from molecular simulations of liquid mixtures.

Keywords: best practices; molecular dynamics simulation; physical property computation

Todo list

I. Preliminaries

Definitions

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- V: Volume
- U: Total energy (including potential and kinetic, excluding external energy such as due to gravity, etc)
- S: Entopy
- N: Number of particles
- T: Temperature
- P: Pressure
- k_B: Boltzmann constant
- β : $(k_B T)^{-1}$
- M: Molar mass
- ρ : Density (M/V)
- *H*: Enthalpy
- G: Gibbs Free Energy (free enthalpy)
- A: Helmholtz Free Energy
- μ : Chemical potential
- *D*: Total dipole moment
- *u*: reduced energy
- f: reduced free energy

Macroscopically, the quantities V,U,N are constants (assuming the system is not perturbed in any way), as we assume that the fluctuations are essentially zero, and any uncertainty comes from our inability to measure that constant value precisely. For a mole of compound (about 18 mL for water), the relative uncertainty in any of these quantites is about 10^{-12} , far lower than any thermodynamics experiment can actually measure.

However, in a molecular simulation, these quantites are not necessarily constant. For example, in an NVT equilib-38 rium simulation, U is allowed to vary. For a long enough simulation (assuming ergodicity, which can pretty much always be assumed with correctly implemented simulations and simple fluids), then the ensemble average value of U = $\langle U \rangle$ will converge to a constant value, and in the limit of large simulations/long time will converge to the macroscopic value U; at least, the macroscopic value of that given model, though perhaps not the U for the real system. In an NVT simulation, clearly V is constant. In an NPT simulation, however, V is a variable, and we must estimate what the macroscopic value would be using the ensemble average $\langle V \rangle$.

The quantities T, P, and μ are typically set as constant during the equilibrium simulations and experiments of interest here. More precisely, the system is in contact with a thermal bath with a fixed T (or in the case of NPT simulations, in contact with a thermal and mechanical bath), and we sample from the systems in equilibrium with this bath. There are a number of quantities that can be used to ESTIMATE constants such as T and P. For example, $\frac{1}{3Nk_B}\sum_i m_i |v_i|^2 >$, where m is the mass of each particle and $|v_i|$ is the magnitude of the velocity of each particle, is an estimate of T (the temperature of the bath), and it's average will be equal to the T. But it is not the temperature. This quantity fluctuates, but T remains constant; otherwise the simulation could not be at constant temperature.

Ensemble averages of some quantity X $(\langle X \rangle)$ are assumed to be averages over the appropriate Boltzmann weighting, i.e. in the NVT ensemble with classical statistical mechanics, they would be $\int X(\vec{x},\vec{p})e^{-\beta U(\vec{x},\vec{p})}d\vec{x}d\vec{p}$. We note that in the limit of very large systems, $\langle X \rangle_{NPT}=\langle X \rangle_{NVT}=\langle X \rangle_{NVT}$.

Ensemble averages can be computed by one of two ways.
First, they can be computed directly, by running a simulation that produces samples with the desired Boltzmann distribution. In that case ensemble averages can be computed as

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 $_{74}$ simple averages, $\langle V \rangle = rac{1}{N} \sum_i V_i$, where the sum is over all $_{111}$ mula for expectations. The free energies can be obtained by $_{75}$ observations. Uncertainties can be estimated in a number of $_{112}$ setting X=1 , and looking at the K equations obtained by different ways, but usually require estimating the number of uncorrelated samples. Secondly, they can be calculated as reweighted estimates from several different simulations, as $_{79}$ $\langle V \rangle = \frac{1}{\sum_i w_i} V_i w_i$ where w_i is a reweighting factor that can be derived from importance sampling theory.

To simplify our discussion of reweighting, we use some additional notation. We define the reduced potential u = $\beta U(\vec{x})$ in the canonical (NVT) ensemble, $u = \beta U + \beta PV$ in the isobaric-isothermal (NPT) ensemble, and $u = \beta U$ – 85 $\beta N\mu$ in the grand canonical ensemble (similar potentials $_{ t 86}$ can be defined in other ensembles). We then define f = $e^{-u}dx$, where the integral is over all of the DOF of the sysss tem (x for NVT, x, V for NPT, and x, N for μVT . For 89 NPT, we then have $f = \beta G$, and for NVT we have f = βA , while for μVT we have $f = -\beta \langle P \rangle V$.

To calculate expectations at one set of parameters generated with paramters that give rise to a different set of proba-93 bility distributions, we start with the definition of an ensemble average given a probability distribution $p_i(x)$.

$$\langle X \rangle_i = \int X(x) p_i(x) dx$$
 (1)

95 We then multiply and divide by $p_i(x)$, to get

$$\langle X \rangle_i = \int X(x) p_i(x) \frac{p_j(x)}{p_j(x)} dx = \int X(x) p_j(x) \frac{p_i(x)}{p_j(x)} dx$$

96 We then note that this last integral can be estimated by the 97 Monte Carlo estimate

$$\langle X \rangle_i = \int X(x) p_j(x) \frac{p_i(x)}{p_j(x)} dx = \frac{1}{N} \sum_{n=1}^N X(x_n) \frac{p_i(x_n)}{p_j(x_n)}$$

 $_{98}$ Where the x_k are sampled from probability distribution 99 $p_i(x)$

butions as: $p_m(x)=rac{1}{N}\sum_{i=1}^N N_k p_k(x)$, where $N=\sum_k N_k$. 140 We can construct a sample from the mixture distribution 141 by simply pooling all the samples from k individual simu- 142 literature sources and the use of reweighting techniques. distribution $p_i(x)$ from samples from the mixture distribu- 144 where suggested by previous authors. 106 tion is:

$$\langle X \rangle_i = \sum_{n=1}^N X(x_n) \frac{p_i(x_n)}{\sum_{k=1}^{N_k} p_k(x_n)} \tag{4}$$

In the case of Boltzmann averages, then $p_i(x) = e^{f_i - u_i(x)}$ where the reduced free energy f is unknown. Reweighting 109 from the mixture distibution becomes.

$$\langle X \rangle_i = \sum_{n=1}^{N} X(x_n) \frac{e^{f_i - u_i(x_n)}}{\sum_{k=1}^{N_k} e^{f_k - u_k(x)}}$$
(5)

which can be seen to be the same formula as the MBAR for-

 $_{113}$ reweighting to the K different distributions.

Finite differences at different temperatures and pressures can be calculated by including states with different reduced potentials. For example, $u_j(x)=\beta_i U(x)+\beta_i(P_i+\Delta P)V$, or $u_j=\frac{1}{k_B(T_i+\Delta T)}U(x)+\frac{1}{k_B(T_i+\Delta T)}P_iV$. However, the relable lationship between f and G can be problematic when looking at differences in free energy with respect to temperature, because $G_2-G_1=eta_2f_2-eta_1f_1$. We can in general write

$$\Delta G_{ij}(T) = k_B T \left(\Delta f_i j(T) - \Delta f_i j(T_{ref}) \right) + \frac{T}{T_{ref}} \Delta G_i j(T_{ref})$$

, where $\Delta G_{ij}(T_{ref})$ is known at some temperature.

Since with MBAR, one can make the differences as small as one would like (you don't have to actually carry out a simulation at those points), we can use the simplest formulas: 125 central difference for first derivatives:

$$\frac{dA}{dx} \approx \frac{1}{2\Delta x} \left(A(x + \Delta x) - A(x - \Delta x) \right)$$

126 And for 2nd deriatives:

$$\frac{d^2A}{dx^2} \approx \frac{1}{\Delta x^2} \left(A(x + \Delta x) - 2A(x) + A(x - \Delta x) \right)$$

127 Thus, only properties at two additional points need to be evaluated to calculate both first and 2nd derivatives.

It may first appear that these finite difference calculations will propagate significant error as they subtract similar numbers. However, MBAR calculates the covariance matrix between $\langle A \rangle$, $A(x+\Delta x)$, and $A(x-\Delta x)$, meaning in practice the uncertainty is far lower than would be expected by standard error propagation of uncorrelated observables.

Note that if the finite differences are re-evaluated using 136 reweighting approaches, it is important that the simulation used generates the correct Boltzmann distribution. If not, 138 reweighted observables will be incorrect, and the results of We now define the mixture distribution of K other distri- 139 the finite difference approach will have significant error.

The following document details calculation of various mechanical observables by both direct methods pulled from lations. The formula for calculating ensemble averages in a 143 Corrections in certain observables are also summarized

Single Phase Properties

Pure Solvent Properties

1. Density

a. Direct calculation Starting with the equation used (5) 149 to calculate the density experimentally,

$$\rho = \frac{M}{V} \tag{6}$$

150 We replace the average with the esemble estimate (calculated either directly, or with reweighting) to obtain:

$$\rho = \frac{M}{\langle V \rangle} \tag{7}$$

b. Derivative Estimate From the differential definition 176 of the Gibbs free energy $dG=VdP-SdT+\sum_i \mu_i dN_i$ that $\eta\eta$ ing these values into the enthalpy equation gives: 154 V can be calculated from the Gibbs free energy as:

$$V = \left(\frac{\partial G}{\partial P}\right)_{TN} \tag{8}$$

155 The density can therefore be estimated from the Gibbs free 156 energy.

$$\rho = \frac{M}{\left(\frac{\partial G}{\partial P}\right)_{T,N}} \tag{9}$$

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157 The derivative can be estimated using a central difference 158 numerical method utilizing Gibbs free energies reweighted 159 to different pressures.

$$\left(\frac{\partial G}{\partial P}\right)_{T,N} pprox \frac{G_{P+\Delta P}-G_{P-\Delta P}}{2\Delta p}$$
 (10)

160 The density can then finally be estimated.

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$$\rho \approx \frac{M}{\frac{G_{P+\Delta P} - G_{P-\Delta P}}{2\Delta P}} \tag{11}$$

This can be calculated from the reduced free energy f if de-162 sired by simply substituting:

$$\rho \approx \frac{\beta M}{\frac{f_{P+\Delta P} - f_{P-\Delta P}}{2\Delta P}} \tag{12}$$

Intuitively, one would imagine that equation 12 would be 184 a worse estimate of density given that the calculations involved have more room for error than direct simulations. 186 That being said, this method should prove invaluable when estimating densities of unsampled states using MBAR.

Molar Enthalpy

This section is on the relation of enthalpy to Gibbs free energy (should we need it). This is not an experimental quan-171 tity, but will be helpful in calculating related properties of interest. The enthalpy, H, can be found from the Gibbs free energy, G, by the Gibbs-Helmholtz relation:

$$H = -T^2 \left(\frac{\partial \left(\frac{G}{T} \right)}{\partial T} \right)_{PN} \tag{13}$$

Transforming the derivative in the Gibbs-Helmholtz rela-175 tion to be in terms of β instead of T yields:

$$H = -T^{2} \frac{\beta^{2}}{\beta^{2}} \left(\frac{\partial \left(\frac{G}{T} \right)}{\partial T} \frac{\partial T}{\partial \beta} \frac{\partial \beta}{\partial T} \right)_{PN} \tag{14}$$

Recall that $\beta = \frac{1}{k_B T}$, therefore $\frac{\partial \beta}{\partial T} = -\frac{1}{k_B T^2}$. Substitut-

$$H = \frac{1}{k_B^3 T^2 \beta^2} \left(\frac{\partial \left(\frac{G}{T} \right)}{\partial \beta} \right)_{P,N}$$

$$= \frac{1}{k_B} \left(\frac{\partial \left(\frac{G}{T} \right)}{\beta} \right)_{P,N} = \frac{\partial f}{\partial \beta}_{P,N} \quad (15)$$

Heat Capacity

The definition of the isobaric heat capacity is:

$$C_P = \left(\frac{\partial H}{\partial T}\right)_{PN} \tag{16}$$

$$C_{P} = \frac{\partial \left(\frac{\partial f}{\partial \beta}\right)}{\partial T}_{P,N} \tag{17}$$

$$C_P = -k_B \beta^2 \frac{\partial^2 f}{\partial \beta^2} \tag{18}$$

This could be computed by finite differences approach or 185 analytical derivation using MBAR

The enthalpy fluctuation formula can also be used to cal-188 culate C_P [?].

$$C_P = \frac{\langle H^2 \rangle - \langle H \rangle^2}{N k_B \langle T \rangle^2} \tag{19}$$

The form is equivalent for isochoric heat capacity, but with derivatives at constant volume rather than pressure.

Isothermal Compressibility

The definition of isothermal compressibility is:

$$\kappa_T = -\frac{1}{V} \left(\frac{\partial V}{\partial P} \right)_T \tag{20}$$

 $_{\mbox{\tiny 195}}$ $\,$ $\,$ a. $\,$ $\,$ $\!$ $\!$ $\!$ $\!$ $\!$ $\!$ $\!$ $\!$ $\!$ Thus, it can be estimated by the fi- $_{\mbox{\tiny 196}}$ nite difference of $\langle V \rangle$

$$\kappa_T = -\frac{1}{2V(T, P)^2} \left(\langle V(P + \Delta P, T) \rangle - \langle V(P - \Delta P, T) \rangle \right)$$

197 Or by the finite differences evaluation of:

$$\kappa_{T} = -\frac{\left(\frac{\partial^{2} G}{\partial P^{2}}\right)_{T,N}}{\left(\frac{\partial G}{\partial P}\right)_{T,N}} = -\frac{\left(\frac{\partial^{2} f}{\partial P^{2}}\right)_{T,N}}{\left(\frac{\partial f}{\partial P}\right)_{T,N}} \tag{22}$$

 κ_T can also be estimated from the ensemble average and fluctuation of volume (in the NPT ensemble) or particle number (in the μ VT ensemble)[?]:

$$\kappa_T = \beta \frac{\langle \Delta V^2 \rangle_{NTP}}{\langle V \rangle_{NTP}} = V \beta \frac{\langle \Delta N^2 \rangle_{VT}}{\langle N \rangle_{VT}}$$
 (23)

5. Speed of Sound

The definition of the speed of sound is[?]:

$$c^{2} = \left(\frac{\partial P}{\partial \rho}\right)_{S} = -\frac{V^{2}}{M} \left(\frac{\partial P}{\partial V}\right)_{S} \tag{24}$$

$$c^{2} = \frac{V^{2}}{\beta M} \left[\frac{\left(\frac{\gamma_{V}}{k_{B}}\right)^{2}}{\frac{C_{V}}{k_{B}}} + \frac{\beta}{V \kappa_{T}} \right]$$
 (25)

Where:

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$$\gamma_V = \left(\frac{\partial P}{\partial T}\right)_V \tag{26}$$

 γ_V is known as the isochoric pressure coefficient. κ_T is the same isothermal compressibility from equation 20

An alternate derivation, applying the triple product rule to $\left(\frac{\partial P}{\partial V}\right)_S$ yields the following.

$$\left(\frac{\partial P}{\partial V}\right)_{S} = \frac{\left(\frac{\partial S}{\partial V}\right)_{P}}{\left(\frac{\partial S}{\partial P}\right)_{V}} \tag{27}$$

$$\left(\frac{\partial S}{\partial V}\right)_{P} = \left(\frac{\partial S}{\partial T}\right)_{P} \left(\frac{\partial T}{\partial V}\right)_{P} = \frac{C_{P}}{T} \left(\frac{\partial T}{\partial V}\right)_{P} = \frac{C_{P}}{TV\alpha} \tag{28}$$

Where $\alpha=\frac{1}{V}\left(\frac{\partial V}{\partial T}\right)_P=\left(\frac{\partial \ln V}{\partial T}\right)_P$ is the coefficient of thermal expansion. The second term in our triple product (22) where $\alpha=\frac{1}{V}\left(\frac{\partial S}{\partial P}\right)_V$, can be expressed as follows:

$$\left(\frac{\partial S}{\partial P}\right)_{V} = \left(\frac{\partial S}{\partial T}\right)_{V} \left(\frac{\partial T}{\partial P}\right)_{V} = \frac{C_{V}}{T} \left(\frac{\partial T}{\partial P}\right)_{V} = \frac{C_{V}}{T\gamma_{V}} \tag{29}$$

Thus our derivation yields:

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$$\left(\frac{\partial P}{\partial V}\right)_{S} = \frac{C_{P}\gamma_{V}}{C_{V}V\alpha} \tag{30}$$

To avoid running an NVT simulation to determine C_V , the following relationship between C_P and C_V can be used:

$$C_P - C_V = TV\left(\frac{\alpha^2}{\kappa_T}\right) \tag{31}$$

This yields the following expression:

$$\left(\frac{\partial P}{\partial V}\right)_{S} = \left(\frac{C_{P}}{C_{P} - TV\frac{\alpha^{2}}{\kappa_{T}}}\right) \left(\frac{\gamma_{V}}{V\alpha}\right) \tag{32}$$

Horn et al set out several ways for calculating α [?]:

 a. Analytical derivative of density with respect to temperature

$$\alpha = -\frac{d\ln\langle\rho\rangle}{dT} \tag{33}$$

b. Numerical derivative of density over range of T of interest The same finite differences approach as shown for isothermal compressibility can be applied here, thus:

$$\alpha = -\frac{d \ln \langle \rho \rangle}{dT} = -\frac{1}{2\rho(T, P)} \left(\ln \langle \rho(P, T + \Delta T) \rangle - \ln \langle V(P, T - \Delta T) \rangle \right)$$
(34)

c. Using the enthalpy-volume fluctuation formula

$$\alpha = \frac{\langle VH \rangle - \langle V \rangle \langle H \rangle}{k_B \langle T \rangle^2 \langle V \rangle} \tag{35}$$

Finite differences approximations and/or analytical derivation can also be used to calculate γ_V or by note of the relation:

$$\gamma_V = -\frac{\alpha}{\kappa_T} \tag{36}$$

Substituting Equations (24),(32), and (36), the following easily calculable expression for speed of sound is obtained:

$$c^{2} = \frac{V}{M\kappa_{T}} \left(\frac{C_{P}}{C_{P} - TV \frac{\alpha^{2}}{\kappa_{T}}} \right) \tag{37}$$

Which can also be written in the simplified form:

$$c^2 = \frac{\gamma}{\rho \kappa_T} \tag{38}$$

Where $\gamma = rac{C_P}{C_V}$.

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6. Dielectric Constant

This equation was provided by a literature reference authored by CJ Fennell[?] and is the standard for calculating the dielectric constant. Below, $\epsilon(0)$ is the zero frequency dielectric constant, V is the system volume and D is the total system dipole moment.

$$\epsilon(0) = 1 + \frac{4\pi}{3k_B T \langle V \rangle} (\langle D^2 \rangle - \langle D \rangle^2) \tag{39}$$

B. Binary Mixture Properties

1. Mass Density, Speed of Sound and Dielectric Constant

The methods for these calculations are the same for a multicomponent system.

2. Activity Coefficient

The definition of chemical potential in a pure substance is:

$$\mu(T,P) = \left(\frac{\partial G}{\partial N}\right)_{T,P} \tag{40}$$

which is a function of only temperaure and pressure.

Then the definition of the chemical potential μ_i of com-

 $_{262}$ pound i in a mixture is:

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$$\mu_i(T, P, \vec{N}) = \left(\frac{\partial G}{\partial N_i}\right)_{T, P, N_i \neq i} \tag{41}$$

 $_{^{264}}$ N_i refers to a molecule of component i and $N_{j \neq i}$ refers to $_{^{265}}$ all molecules other than component i, with \vec{N} the vector of all component numbers. Since μ_i is intensive, this is equivalently a function of the vector of mole fractions \vec{x}_i instead of simply of N_i .

For an ideal solution, the chemical potential μ_i can be re- μ_i lated to the pure chemical potential by

$$\mu_i(T, P, \vec{x}_i) = \mu(T, P) + k_B T \ln{(\gamma_i)}$$
 (42)

By analogy to this form, we can say

$$\mu_i(T, P, \vec{x}_i) = \mu(T, P) + k_B T \ln(x_i \gamma_i)$$
(43)

²⁷⁷ Where γ_i is the activity coefficient of component i, and i is a function of T,P,and \vec{x}_i . Rearrangement of the previous equation yields:

$$\gamma_i = \frac{e^{\left(\frac{\mu_i(T,P,\vec{x}_i) - \mu(T,P)}{k_B T}\right)}}{x_i} \tag{44}$$

Although chemical potentials cannot be directly calcu- lated from simulation, chemical potential residuals can. We (39) lated from simulation, chemical potential residuals can. We (39) can calculate the difference $\mu_i(T,P,\vec{x}_i)-\mu(T,P)$ by calculating $\Delta\mu(T,P)_{liquid}-\Delta\mu(T,P)_{gas}$ using a standard alchemical simulation of the pure substance, followed by the calculation of $\mu_i(T,P,\vec{x}_i)_{liquid}-\Delta\mu(T,P,\vec{x}_i)_{gas}$, and assuming that $\Delta\mu(T,P,\vec{x}_i)_{gas}=\Delta\mu(T,P)_{gas}$. Note: there are a few subleties here relating to the $\ln x_i$ factor, but it appears that with alchemical simulations with only one particle that is allowed to change, this will cancel out (need to follow up).

Several of these alchemical simulation methods for calculating activity coefficients have been pioneered by Andrew Paluch [?]. A method detailing the calculation of infinite dilution activity coefficients γ_i^{inf} for binary a mixture follows directly:

$$\begin{split} \ln \gamma_2^{\infty}\left(T,P,x_2=0\right) &= \beta \mu_2^{res,\infty}\left(T,P,N_1,N_2=1\right) \\ &+ \ln \left[\frac{RT}{V_1\left(T,P\right)}\right] - \ln f_2^0\left(T,P\right) \end{split} \tag{45}$$

Where $\beta\mu_2^{res,\infty}$ is the dimensionless residual chemical

potential of component 2 at inifinite dilution. The residual is 343 302 defined here as the difference between the liquid and ideal gas state. $V_1(T, P)$ is the molar volume of component 1 at T and P. $\ln f_2^0(T,P)$ is the natural logarithm of the pure $_{305}$ liquid fugacity of component 2 and is defined as:

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$$\ln f_2^0(T, P) = \beta \mu_2^{res}(T, P) + \ln \left[\frac{RT}{V_2(T, P)} \right]$$
 (46)

Paluch et al. use a multistage free energy perturbation approach utilizing MBAR in order to calculate the residual 350 partial molar Gibbs free energy and dimensionless Gibbs free energy differences between multiple states are readily computed with MBAR). The idea is to connect two states of interest. In the case of a pure liquid, connecting a system of pure liquid molecules with N-1 interacting molecules and one fully decoupled molecule to a system of N fully interacting molecules. The coupling/decoupling process is detailed by Paluch et al [?], but involves a linear alchemical switching function where LJ and electronic interactions are 320 slowly turned on for the decoupled molecule until they are 355 ³²¹ fully on. The free energy of this coupling is calculated by simpling summing the free energy changes along this path.

3. Excess Molar Properties

The general definition of an excess molar property can be 325 stated as follows:

$$y^E = y^M - \sum_i x_i y_i \tag{47}$$

Where y^E is the excess molar quantity, y^M is the mixture $\,^{\scriptscriptstyle 360}$ quantity, x_i is the mole fraction of component i in the mixture and y_i is the pure solvent quantity. In general, the simplest methods for calculating excess molar properties for binary mixtures will require three simulations. One simulation is run for each pure component and a third will be run for the specific mixture of interest. We note that only one set of pure simulations are needed to calculate excess properties at all compositions.

Excess Molar Heat Capacity and Volume

Excess molar heat capacities and volume will be calculated using the methods for the pure quantities in section Iin combination with the general method for excess property calculation above.

Excess Molar Enthalpy

Excess molar enthalpy can be calculated using the general relation of molar enthalpy as it relates to Gibbs Free En- $_{346}$ ergy from section I and the general method of excess molar property calculation above or by the following[?]:

$$H^{E} = \langle E^{M} \rangle + PV^{E} - \sum_{i} x_{i} \langle E_{i} \rangle \tag{48}$$

Where $\langle E \rangle$ denotes an ensemble average of total energy chemical potentials (recall that the chemical potential is the 351 and V^E is calculated using the general method of excess 352 molar properties.

Suggested Corrections

Heat Capacity

Horn et al suggest a number of vibrational corrections be applied to the calculation of C_P due to a number of approx-358 imations made during the simulation of the liquid [?]. The 359 following terms were added as a correction.

$$\left(\frac{\partial E_{vib,l}}{\partial T}\right)_{P} = \left(\frac{\partial E_{vib,l,intra}^{QM}}{\partial T}\right)_{P} + \left(\frac{\partial E_{vib,l,inter}^{QM}}{\partial T}\right)_{P} - \left(\frac{\partial E_{vib,l,inter}^{CM}}{\partial T}\right)_{P} \tag{49}$$

Where:

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$$\left(\frac{\partial E_{vib}^{CM}}{\partial T}\right)_{P} = k_{B} n_{vib} \tag{50}$$

$$\left(\frac{\partial E_{vib}^{QM}}{\partial T}\right)_{P} = \sum_{i=1}^{n_{vib}} \left(\frac{h^{2}v_{i}^{2}e^{\frac{hv_{i}}{k_{B}T}}}{k_{B}T^{2}\left(e^{\frac{hv_{i}}{k_{B}T}} - 1\right)^{2}}\right)$$
(51)

Above, n_{vib} is the number of vibrational modes, h is $_{ exttt{368}}$ Planck's constant and v_i is the vibrational frequency of $_{369}$ mode i.

III. Properties Involving Change of Phase

A. Pure Solvent Properties

1. Enthalpy of Vaporization

The definition of the enthalpy of vaporization is[?]:

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$$\Delta H_{vap} = H_{gas} - H_{liq} = E_{gas} - E_{liq} + P(V_{gas} - V_{liq})$$
 (52)

If we assume that $V_{gas}>>V_{liq}$ and that the gas is ideal (and therefore kinetic energy terms cancel):

$$\Delta H_{vap} = E_{gas,potential} - E_{liq,potential} + RT$$
 (53)

B. Suggested Corrections

1. Enthalpy of Vaporization

An alternate, but similar, method for calculating the enthaps lapy of vaporization is recommended by Horn et al [?].

$$\Delta H_{vap} = -\frac{E_{liq,potential}}{N} + RT - PV_{liq} + C$$
 (54)

In the above equation ${\cal C}$ is a correction factor for vibrational energies, polarizability, non-ideality of the gas and pressure. It can be calculated as follows.

$$C_{vib} = C_{vib,intra} + C_{vib,inter}$$

$$= (E_{vib,QM,gas,intra} - E_{vib,QM,liq,intra})$$

$$+ (E_{vib,QM,liq,inter} - E_{vib,CM,liq,inter})$$
(55)

The QM and CM subscripts stand for quantum and classical mechanics, resectively.

$$C_{pol} = \frac{N}{2} \frac{(d_{gas} - d_{liq})^2}{\alpha_{n,gas}} \tag{56}$$

Where d_i is the dipole moment of a molecule in phase i and $\alpha_{p,gas}$ is the mean polarizability of a molecule in the gas phase.

$$C_{ni} = P_{vap} \left(B - T \frac{dB}{dT} \right) \tag{57}$$

Where B is the second virial coefficient.

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$$C_{x} = \int_{P_{ext}}^{P_{vap}} \left[V\left(P_{ext}\right) \left[1 - \left(P - P_{ext}\right) \kappa_{T} \right] - TV\alpha \right] dP$$
(58)

Where P_{ext} is the external pressure and $V\left(P_{ext}\right)$ is the volume at P_{ext} .

This is frequently done as a single simulation calculation by assuming the average intramolecular energy remains constant during the phase change, which is rigorously correct for something like a rigid water molecule (intramolecular energies are zero), but less true for something with structural rearrangement between gas and liquid phases.

As discussed by myself and MRS, we have decided to not initially begin the parametrization process using enthalpy of vaporization data. While force field parametrization is commonly done using said property we have ample reason to not follow classical practice. First of all, the enthalpy data is usually not collected at standard temperature and pressure, but at the saturation conditions of the liquid being vaporized [?]. This would require corrections to be made to get the property at STP (the process will be explained below) using fitted equations for heat capacity. Not only is this inconvenient, but it adds an unknown complexity when adjusting experimental uncertainties due to the added correction. Often times the uncertainties of these "experimental" enthalpies are unrecorded because they are estimated from fitted Antoine equation coefficients [?].

An additional issue is the necessity of having to use gas
phase simulation data in order to validate a parametrization process meant for small organic liquids and their mixtures. Following an example of Wang et al. [?] we plan to
mixed use enthalpy of vaporization calculations as an unbiased means of testing the success of the parametrization. If
the parametrization procedure is expanded to use enthalpy
of vaporization, corrections can be made to the experimental data in order to get a value at STP using the following
equation.

$$\Delta H_{vap}(T) = \Delta H_{vap}^{ref} + \int_{T_{ref}}^{T} \left(C_{P,gas} - C_{P,liq} \right) dT \quad (59)$$

IV. Numerical Considerations

A. Quality of Timeseries Data

With the finite computing resources that are available to- 438 day, it is often necessary to make trade-offs between the 439 uncertainty of the result and the amount of time/resources

440 spent. As a major assumption of Molecular Dynamics simu- 495 period, samples close to each other in time are likely corcific problem being addressed?

of the simulation, often measured in time units such as 500 pling is used for all calculations. nanoseconds or microseconds, and the size of the system that is simulated, measured in number of molecules. Because timescales vary based on the nature of the system, a more general metric that we will use in this discussion is N, the number of uncorrelated equilibrium samples. We will denote the number of molecules as M in this discussion to distinguish the two.

1. Equilibrium Sampling

The relationship between N and uncertainty is straightforward; as one collects more samples of the same system, the standard deviation decreases as $\mathcal{O}(N^{-\frac{1}{2}})$, as would be expected for the standard formula for sample standard deiation. It is worth noting that while N follows this rule, the length of the simulation will not follow it exactly. This is because the detection of equilibration and the choice of uncorrelated equilibrium samples will vary between users and

For a timeseries to satisfy the Ergodic Hypothesis and allow for calculation of equilibrium samples, the system needs to have settled into an equilibrium region, following a "burn-in" period in which the system starts at some nonequilibrium configuration and relaxes into an equilibrium configuration. This portion of a timeseries needs to be removed; the question as to how to do so is slightly trickier. For users running many simulations of similar systems, often a burn-in time is determined by inspection and then set for all subsequent simulations. However, this can backfire based on the simulation parameters being varied. For example, systems with small numbers of molecules can equilibrate much slower than larger systems, making the choice of equilibrium for one invalid for another, or discarding valuable equilibrium data. Another option is an automatic equilibration detection algorithm, like the one proposed by Chodera. In this work, this automatic equilibration algo- 527 rithm is used in all analysis. It is important to note, however, that algorithms like this should not be used in a vacuum; the user should periodically check to make sure that the algorithm is functioning properly, and have a rudimentary understanding of how it works.

494 system may be at equilibrium constantly after the burn-in 532 This relationship is shown in figure:

lations is the Ergodic Hypothesis, sampling over a timeseries 496 related. Two common methods for obtaining uncorrelated will only be equivalent in the limit of infinite time. In all other 497 samples are through subsampling and block averaging. In cases, the timeseries generated from MD will function as an 498 block averaging, the timeseries is separated into "blocks" approximation to phase space sampling. As this is the case, 499 of data and observations are taken from the mean of each there is an inherent decision to be made in the collection of 500 block. This requires for a size of block to be chosen, and as all timeseries data: how much effort does one need to spend 501 the block size increases, the variance of the observable will to get a result with the appropriate confidence for the spe- 502 eventually reach a plateau where samples are uncorrelated. 503 Another method is subsampling, where the statistical ineffi-For bulk thermophysical properties, there are two major $_{504}$ ciency g is estimated and then snapshots are chosen based factors that consume computational resources: the length 505 on this parameter, without averaging. In this work, subsam-

System Size

For the number of molecules M, relationships with uncertainty are more complicated and vary from property to prop- $_{510}$ erty. This can lead to unintuitive results in which a larger Mmay reduce uncertainty in one physical quantity, but for a derived property of said quantity, the same increase in Mmay not improve estimates.

Enthalpy and Heat Capacity

This is best illustrated with the example of energy and 516 heat capacity. From Eq. 19, the extensive heat capacity is directly proportional to the variance in the enthalpy, which 518 is defined as $\langle H^2 \rangle - \langle H \rangle^2$. For any extensive property, its value has a linear relationship with M; as you go from 100 molecules to 200 molecules, the total heat capacity (or enthalpy, or volume, etc.) will double.

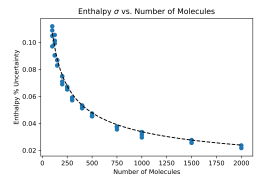
Therefore, the heat capacity goes as $C_P \sim \mathcal{O}(M)$, 523 and subsequently, so does the variance of the enthalpy: $Var(H) \sim C_P \sim \mathcal{O}(M)$. If we wish to examine the variance of the molar enthalpy, an intensive quantity, we can make use of this standard formula for variances:

$$Var(aX) = a^2 Var(X) \tag{60}$$

Since molar enthalpy is defined as $h = \left(\frac{H}{M}\right)$, the variance 528 of the molar enthalpy is given by:

$$Var(h) = Var\left(\frac{H}{M}\right) = \frac{1}{M^2}Var(H) \sim \frac{\mathcal{O}(M)}{M^2} \sim \mathcal{O}(1/M)$$
(61)

Another important factor in the calculation of thermo- 529 So, the standard deviation of the molar enthalpy goes as physical properties is the collection of uncorrelated equilib- $_{530}$ $\mathcal{O}(M^{-\frac{1}{2}})$, meaning that quadrupling the size of the system rium samples from an equilibrated timeseries. Although the $_{531}$ will cut the uncertainty in half, just as quadrupling N would.



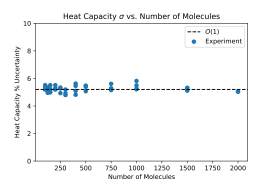


FIG. 1: Figure shows enthalpy % uncertainty (calculated as $(\sigma_h/h) \times 100\%$) FIG. 2: Figure shows heat capacity % uncertainty (calculated as vs. M for simulations of liquid cyclohexane at 293.15 K, 1.01 bar, calculated (σ_{C_P}/C_P) imes 100%) vs. M for simulations of liquid cyclohexane at 293.15 K, 1.01 bar, calculated via fluctuations. No dependence on M. via fluctuations. $\mathcal{O}(M^{-\frac{1}{2}})$ on M.

For the isothermal compressibility, the variance is easy to Now, we examine the uncertainty behavior of C_P , the examine the variance is easy to Now, we examine the uncertainty behavior of C_P , the example of the variance of the volume is known. From tensive heat capacity. Since we know $C_P \sim Var(H)_{i,0}^{554}$ tensive heat capacity. Since we know $C_P \sim Var(H)_{i,0}^{554}$ teq. 23, we can see that with constant temperature, the variance of the variance of the variance of κ_T is as such: tributed samples, the variance of the variance is given by:

$$Var(Var(X)) \approx \frac{2(N-1)}{N^2} (Var(X))^2$$
 (62) $Var(\kappa_T) \sim Var\left(\frac{Var(V)}{V}\right) = \frac{1}{V^2} Var(Var(V))$ (65)

Holding N constant and applying to the enthalpy, we obtain the following:

$$Var(\kappa_T) \sim \frac{1}{V^2} (Var(V))^2 \sim \frac{\mathcal{O}(M^2)}{\mathcal{O}(M^2)} \sim \mathcal{O}(1)$$
 (66)

$$Var(C_P) \sim Var(Var(H)) \sim (Var(H))^2 \sim \mathcal{O}(M^2)$$
 (63) the r

So, the uncertainty of the compressibility is independent of (63) the number of molecules used to simulate the system. This ses claim is supported by experimental data in figure 3:

As we examine the *molar* heat capacity $c_P = C_P/M$ and ₅₄₀ apply the variance formula from above, we obtain:

$$Var(c_P) = Var(\frac{C_P}{M}) = \frac{Var(C_P)}{M^2} \sim \frac{\mathcal{O}(M^2)}{M^2} \sim \mathcal{O}(1)$$
 (64)

So, even though estimates of the molar enthalpy h improve as the size of the system increases, estimates of the molar heat capacity c_P , which is alternatively defined as the derivative of h with respect to T, do not. Note that while this derivation is for NPT systems, a similar derivation follows for NVT systems, replacing H with E and C_P with C_V .

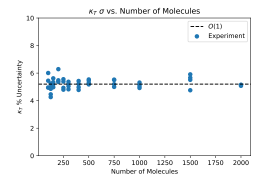


FIG. 3: Figure shows speed of sound % uncertainty (calculated as $(\sigma_{\kappa_T}/\kappa_T) \times 100\%$) vs. M for simulations of liquid cyclohexane at 293.15 K, 1.01 bar, calculated via fluctuations. No dependence on M.

Volume and compressibility

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For the volume V, the standard deviation also goes as $\mathcal{O}(M^{-\frac{1}{2}})$. This follows from the information we already know about energy and enthalpy and the relationship 5%tween the two. Since they are related by H=E+PV, and the variance of H and E both go as $\mathcal{O}(M)$ due to their rela-

ported by experimental data in figure.

Speed of Sound

For thermophysical properties that are calculated from tionship with the heat capacities C_P and C_V , the variance of several other basic properties, such as speed of sound c=V must also go as $\mathcal{O}(M)$. By similiar analysis to that for the $rac{\gamma}{
ho\kappa_T}$, it is important to examine the relative contributions of molar enthalpy h, it follows that the standard deviation \mathfrak{softhe} error to determine the overall uncertainty behavior of the molar volume v goes as $\mathcal{O}(M^{-\frac{1}{2}}$, and this claim is sup-the property. For example, γ and κ_T are both properties with uncertainties that do not depend on M, but ρ should

have an $\mathcal{O}(N^{-\frac{1}{2}})$ uncertainty behavior, similar to molar vol- $_{608}$ these all methods are equivalent, or if there are differences behavior to be dependent on M, but data from experiment $_{610}$ depending on the situation. 576 in Figure 4 shows no correlation.

ume. Therefore, one would expect the overall uncertainty 609 in methods which may make them more or less attractive,

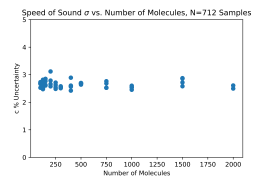


FIG. 4: Figure shows speed of sound % uncertainty (calculated as (σ_c/c) × 100%) vs. M for simulations of liquid cyclohexane at 293.15 K, 1.01 bar, calculated via fluctuations. No apparent dependence on M.

Discussion

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From these analyses, we can see that properties taken from direct averages of timeseries data, (V, E, H) have standard deviation that goes as $\mathcal{O}(M^{\frac{1}{2}})$. The molar versions of these properties subsequently have standard deviation that goes as $\mathcal{O}(M^{-\frac{1}{2}})$. However, this does not imply the same for properties calculated via fluctuations. Fluctuation properties can have standard deviation as $\mathcal{O}(M)$, like C_P , or with no dependence on M, like c_P or κ_T . Additionally, they can have a very weak dependence on M, such as speed of sound c. Because of these varying dependencies between quantities, and the extra computational cost of a larger system size, we recommend that additional computational resources be used to collect more equilibrium samples, which has a consistent effect on uncertainty for all properties. This recommendation comes with the caveat that systems need to be large enough to avoid finite size effects. When systems are too small, systems may not equilibrate very quickly, amongst other problems. For a more in-depth discussion of finite size effects, see. It is also important to remember when modifying the number of molecules n a system that Molecular Dynamics programs report energies in units of energy/mole of box, rather than energy per mole of compound. For pure compounds, one must divide the "moles of box" value by the number of molecules in order to get a true molar energy.

Comparison of Calculation Methods

₆₀₇ properties. Therefore, it is worth investigating whether ₆₅₈ is more like 75% for κ_T .

Comparison of MBAR finite differences and direct finite differences

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Suppose we are interested in calculating a thermodynamic property B at state X, defined as B = dA/dX. When calculating this property from timeseries data via centered finite difference (such as in Eq. 10,21), the simplest and most nature choice is to choose a finite difference ΔX , perform simulations at $X - \Delta X$ and $X + \Delta X$, take the expectations $\langle A(X-\Delta X)\rangle$ and $\langle A(X+\Delta X)\rangle$, and then calculate the finite difference from these two values ("Direct Fi-621 nite Differences"). Another possibility is take a simulation $_{622}$ at X, and use reweighting via the MBAR method [?] to estimate A at $X - \Delta X$ and $X + \Delta X$, then calculate the finite difference from these estimates ("MBAR Finite Differences"). The choice of finite difference is important here, as ΔX will determine the accuracy of the calculation. Theoretically, we would like the smallest finite difference possi-₆₂₈ ble to minimize potential bias in the calculation. For direct finite differences, however, a small ΔX will provide unreliable estimates, as timeseries data at very similar state points are highly correlated, leading to high uncertainty. As ΔX increases, the timeseries data becomes less correlated and uncertainty goes down, but there is potential for bias, which 634 depends on the behavior of the function.

When using MBAR finite differences, the problem is opposite: MBAR estimates are inherently decorrelated, leading to decent uncertainty at small ΔX , but MBAR requires significant phase overlap between the simulated state and the desired state, so estimates become more unreliable as $_{640}$ ΔX increases. However, since MBAR finite differences are stable over a large range of $\Delta X/X$ ($(10^{-6}-10^{-3})$ for heat 642 capacities), this is less of an issue. However, it does make comparisons of direct finite differences and MBAR finite differences difficult, as the minimum ΔX in direct finite differences that produces uncertainties equivalent to MBAR finite differences is roughly 10^{-2} in heat capacity; MBAR estimates are already poor at this point. In order to quantify the relia-648 bility of the MBAR finite difference, the number of effective $_{ ilde{6} ext{49}}$ samples (N_{eff}) metric can be used. This metric quantifies 650 how many of the samples from simulated state A could be 651 drawn from reweighted state B. Previous use of this metric $_{
m 652}$ has shown that a rough minimum of $N_{eff}=50$ is required 653 for statistics to be stable, but for accurate finite differences, the percentage of effective samples $((N_{eff}/N) \times 100\%)$ is For the properties discussed in this paper, the methods of 655 a better metric, but may depend on the property being calcalculation generally fall into one of two categories: deriva- $_{656}$ culated. For example, MBAR finite differences for C_P agree tives via finite differences or calculations from fluctuation 657 with fluctuations with roughly 10 % of effective samples, but

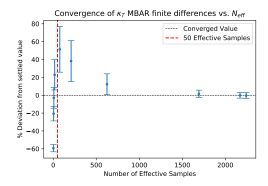


FIG. 5: Percent deviation from fluctuation value as a function of N_eff . Dashed red line indicates minimum requirement of 50 samples. Error bars indicate bootstrapped 95% CI

Since the number of effective samples is easily adjusted, as a best practice, we recommend a percentage of effective samples of 90 % or higher. It is also important to recognize that for properties that require reweighting in multiple variables, such as speed of sound, the number of effective samples for all variables needs to be taken into account (in the example of speed of sound, T and P). In general, comparison with fluctuation calculations is the ultimate metric for accuracy of MBAR finite differences.

To test this, we performed a 10 ns simulation of cyclohexane at 293.15 K and 1.01 bar, as well as several pairs of 5 ns cyclohexane simulations, also at 1.01 bar, but spaced at intervals of 0.5, 1, 2, 5, 7.5, and 10 K apart from 293.15 K. Heat capacity calculations from finite differences were performed via MBAR on the 10 ns simulation, and directly on the pairs of 5 ns simulations.

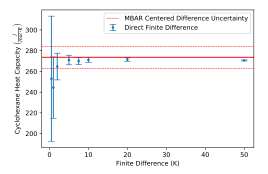


FIG. 6: Comparison of direct finite difference methods and MBAR reweighted finite differences for liquid cyclohexane C_P at 293.15 K, 1.01 bar. Red lines show MBAR estimate and uncertainty, error bars indicate bootstrapped 95 % CI.

2. Finite differences and fluctuations

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For properties that can be calculated via finite differences or fluctuations, such as κ_T (Eq. 21, 23), C_P (Eq. 16, 19), or α_R (Eq. 33,35), the two methods should be theoretically equivablent, as fluctuation formulas are derived by substituting sta-

tistical mechanical expressions for energy, enthalpy, or volume into derivative formulas and evaluating the derivative. In the limit of infinitely small finite differences and infinitely long timeseries, this will be exactly true; however, since neither condition can be met, it is important to understand if these methods are approximately equivalent under normal conditions.

Because of the large number of particles and the chaotic nature of the interactions between them, replicate timeseries are often slightly different, and will provide substantially different sets of uncorrelated equilibrium samples. Therefore, agreement between finite differences and fluctuation methods are not based on agreement to some "grand truth" but agreement between the data used to compute the derivative and the fluctuations. This point is illustrated with the following relation:

$$\frac{H(T_2) - H(T_1)}{T_2 - T_1} \approx \left(\frac{\partial H}{\partial T}\right)_P = \left(\frac{Var(H)}{RT^2}\right)_{\infty} \approx \left(\frac{Var(H)}{RT^2}\right)$$
(67)

As fluctuation calculations and finite differences are both approximations to the "grand truth" of true derivatives and true fluctuations, they will not agree unless they come from the same data source. For direct finite differences, since simulations are taken at different conditions than the simulation used for the fluctuation calculation $(X+\Delta X)$ and $X-\Delta X$ instead of X, the agreement between fluctuations and finite differences is significant. In the case of MBAR finite differences, the same data can be used to calculate the finite differences and the fluctuations, so these two methods should agree, up to numerical error caused by the finite difference size chosen for MBAR. In this way, MBAR finite differences and fluctuations are a good sanity check, as they should be almost identical if implemented correctly.