

Free energy calculations

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Why do free energy calculations?

The free energy G gives the population of states:

$$\frac{P_1}{P_2} = \exp\left(\frac{\Delta G}{k_B T}\right) , \quad \Delta G = G_2 - G_1$$

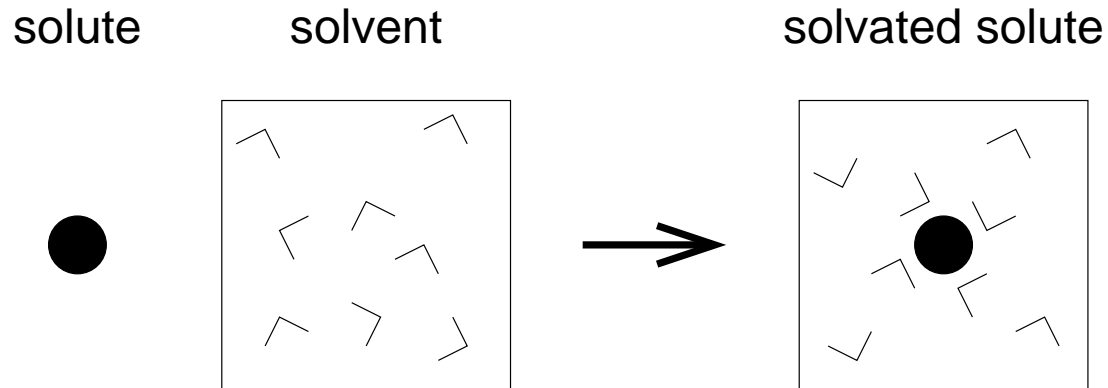
Since we mostly simulate in the NPT ensemble we will use the Gibbs free energy G (not the NVT Helmholtz free energy A)

A free energy difference can be split into two terms:

$$\Delta G = \Delta H - T\Delta S , \quad \Delta H = \Delta U + P\Delta V$$

ΔG is easier to determine than ΔU

Free energy of solvation



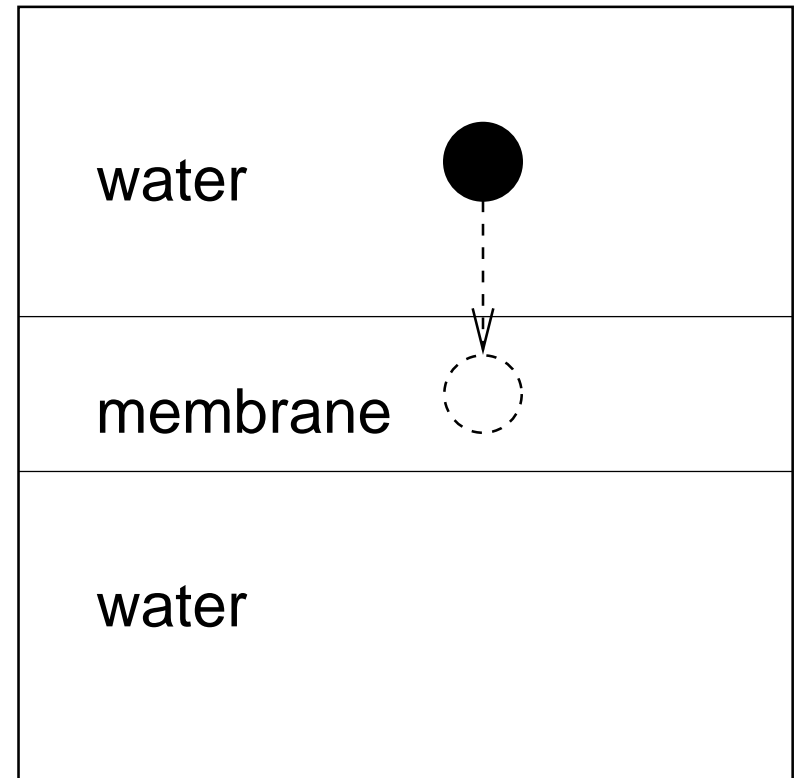
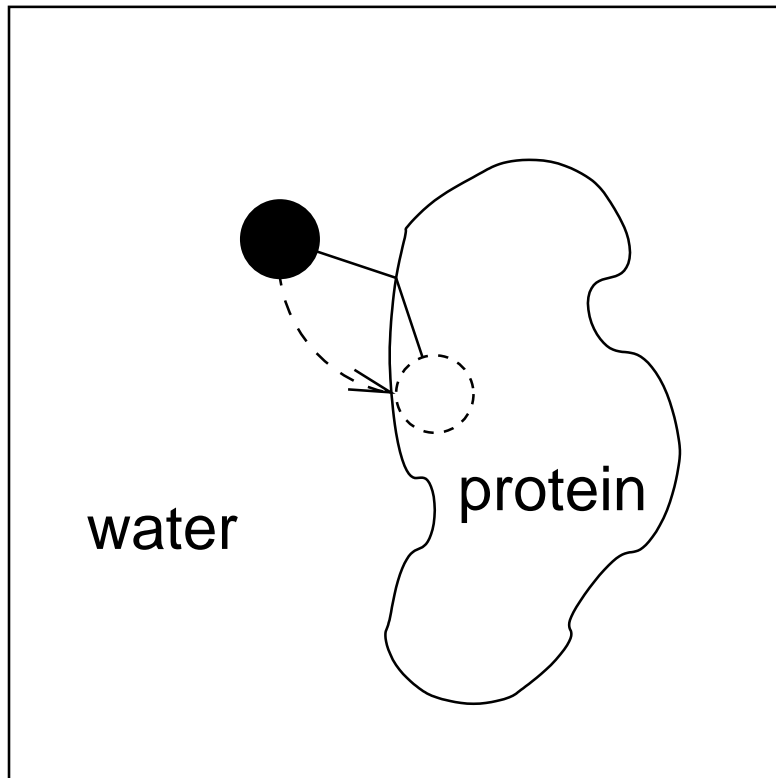
ΔG of solvation is often used to parametrize force fields

Partitioning properties:

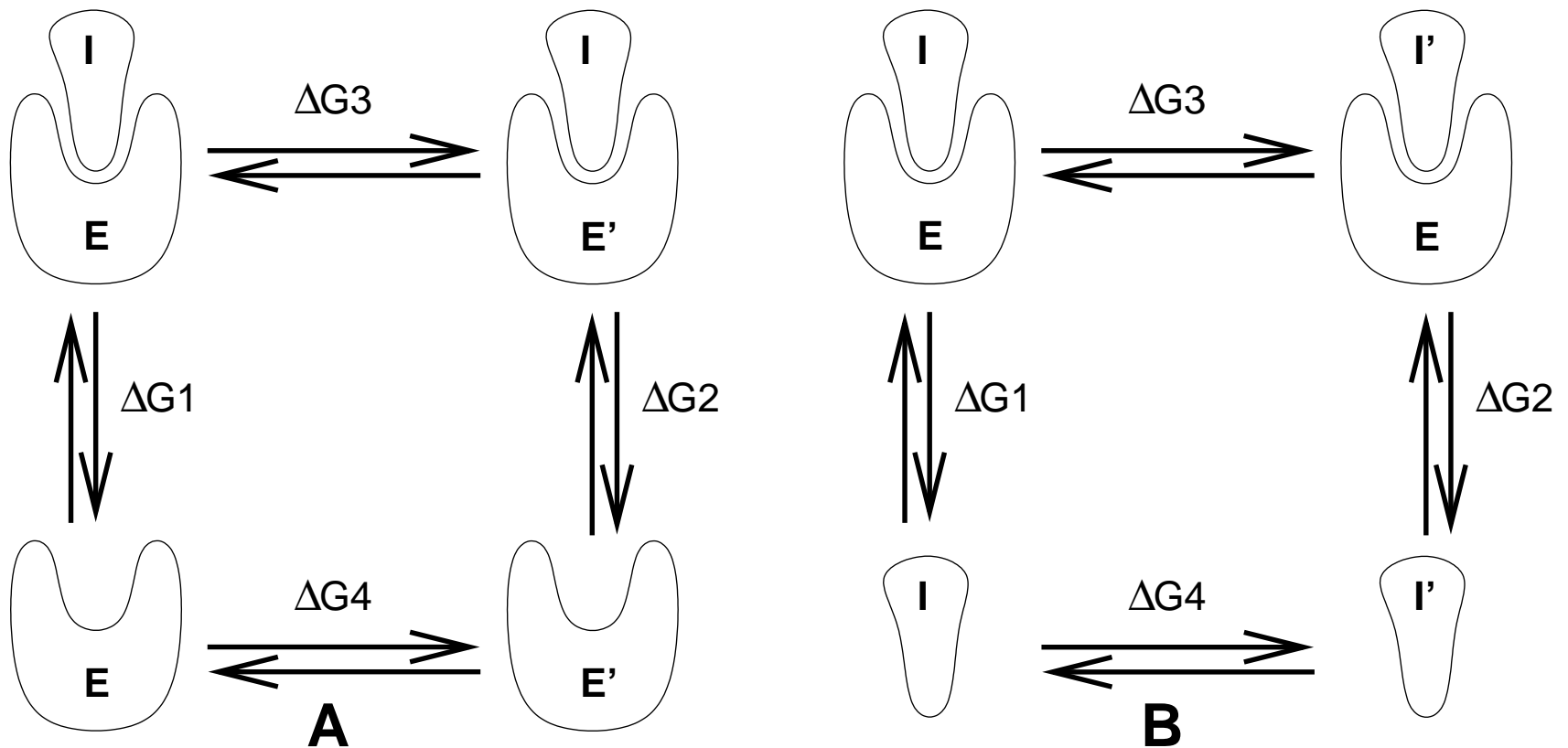
similarly one can determine the free energy of transfer from a polar (water) to an apolar solvent (octane, cyclohexane)

This is important for protein folding and peptide-membrane interactions

Partitioning free-energies

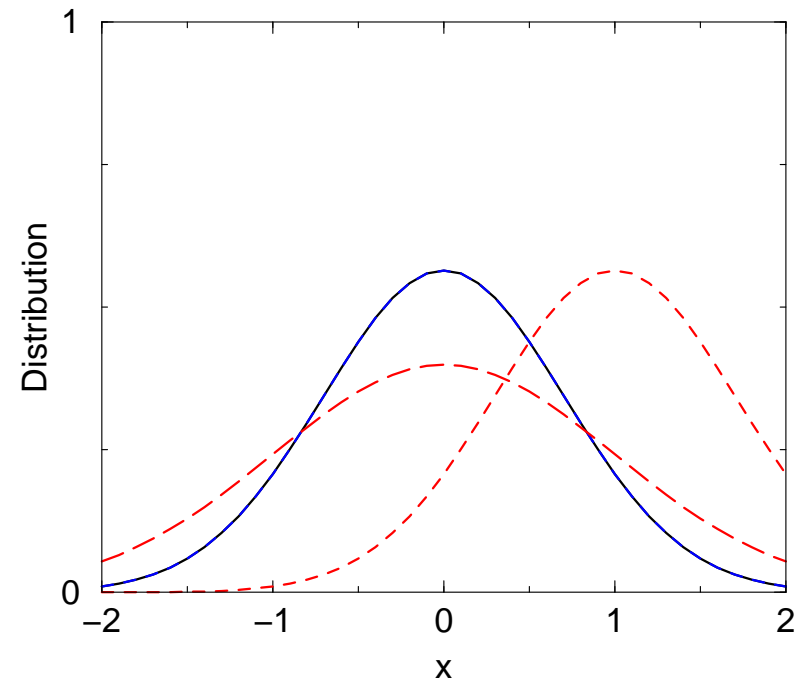
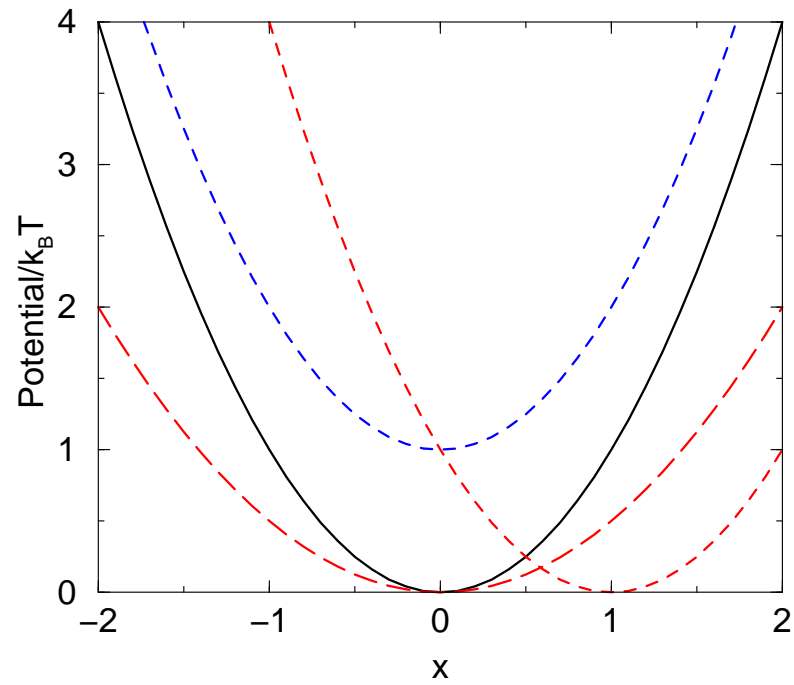


Binding free energies



$$\Delta\Delta G = \Delta G_1 - \Delta G_2 = \Delta G_3 - \Delta G_4$$

Distributions



ΔG between similar states

When the states of interest are very similar, e.g. small changes in charge or LJ parameters, only the potential energy difference matters and one can do single step perturbation

In Gromacs this can be done in two ways:

- Run a simulation for state A
- Do `mdrun -rerun traj.trr` with B-state parameters
- Determine the potential energy difference

or

- Run a simulation for state A with also a B-state topology and get the potential energy difference from the free energy code

ΔG between different states

With frequent transitions between states and small ΔG :

- simulate the system
- count the populations

With infrequent transitions between states or large ΔG :

We can modify the Hamiltonian to gradually move the system from state A to state B

The free energy difference is then given by the total work associated with changing the Hamiltonian

The coupling parameter approach

We add a coupling parameter λ to the Hamiltonian:

$$H = H(p, q; \lambda)$$

$$H(p, q; 0) = H^A(p, q) \quad , \quad H(p, q; 1) = H^B(p, q)$$

The free energy difference between state A and B is then given by:

$$G^B(p, T) - G^A(p, T) = \int_0^1 \left\langle \frac{\partial H}{\partial \lambda} \right\rangle_{NPT; \lambda} d\lambda$$

The coupling parameter approach

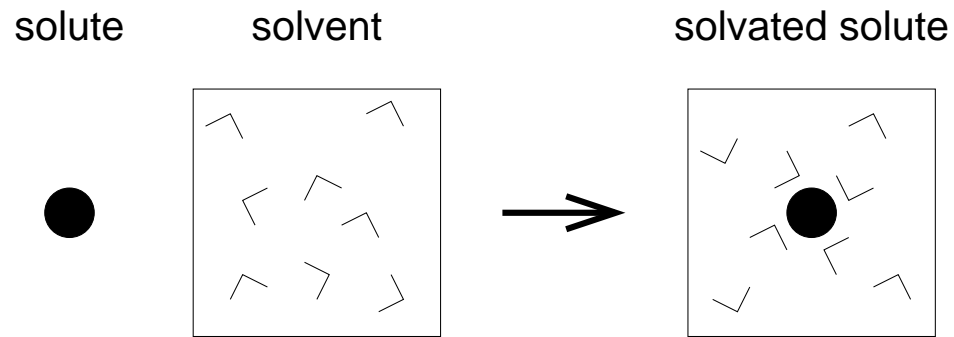
The λ dependence of $H(p, q; \lambda)$ can be chosen freely, as long as the end points match the states

The simplest approach is linear interpolation:

$$H(p, q; \lambda) = (1 - \lambda)H^A(p, q) + \lambda H^B(p, q)$$

This works fine, except when potentials with singularities are affected
(LJ, Coulomb)

The coupling parameter approach



State A: solute fully coupled to the solvent

State B: solute fully decoupled from the solvent

An example approach:

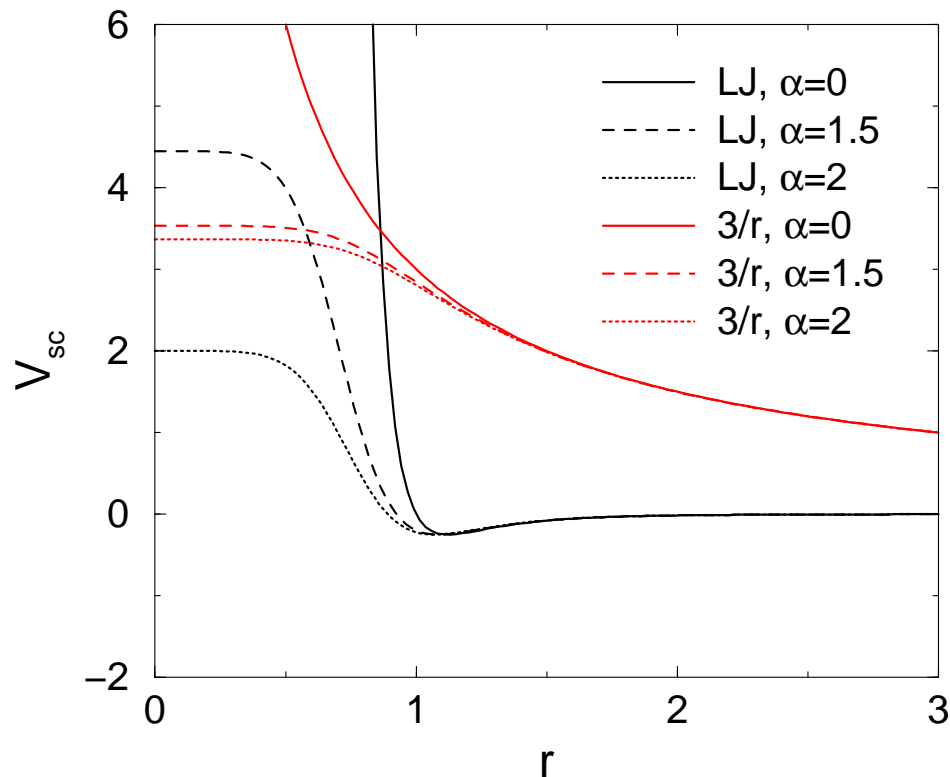
Run simulations at: $\lambda = 0, 0.1, \dots, 0.9, 1.0$

Soft-core interactions

Instead of linear interpolation we use:

$$V_{sc}(r) = (1 - \lambda)V^A(r_A) + \lambda V^B(r_B)$$

$$r_A = (\alpha\sigma_A^6\lambda^p + r^6)^{\frac{1}{6}}, \quad r_B = (\alpha\sigma_B^6(1 - \lambda)^p + r^6)^{\frac{1}{6}}$$



Solvation free energy in practice

Make a topology where:

- the B-state atom types of the solute have LJ parameters zero
- the B-state charges of the solute are zero

1) Simulate the system with solvent at several λ -values 0, ..., 1

2) Simulate the system in vacuum at several λ -values 0, ..., 1

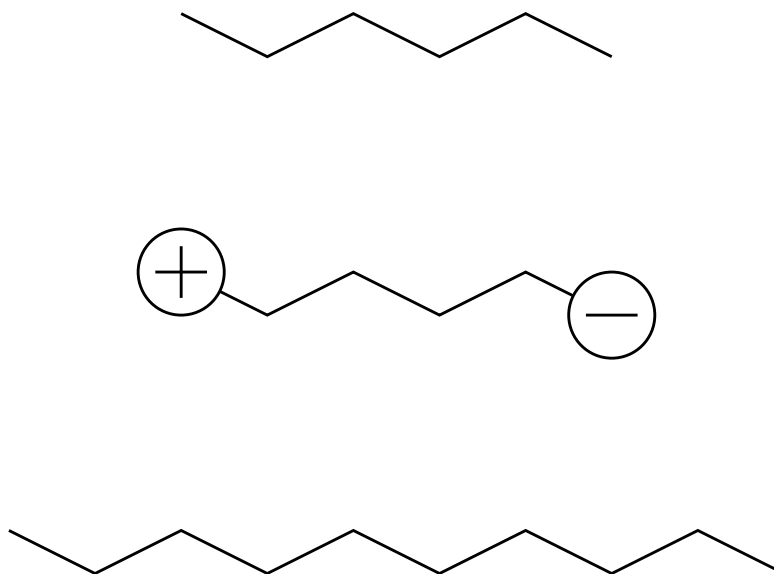
The solvation free energy is given by:

$$\Delta G_{solv} = \int_0^1 \left\langle \frac{\partial H}{\partial \lambda} \right\rangle_{2)} d\lambda - \int_0^1 \left\langle \frac{\partial H}{\partial \lambda} \right\rangle_{1)} d\lambda$$

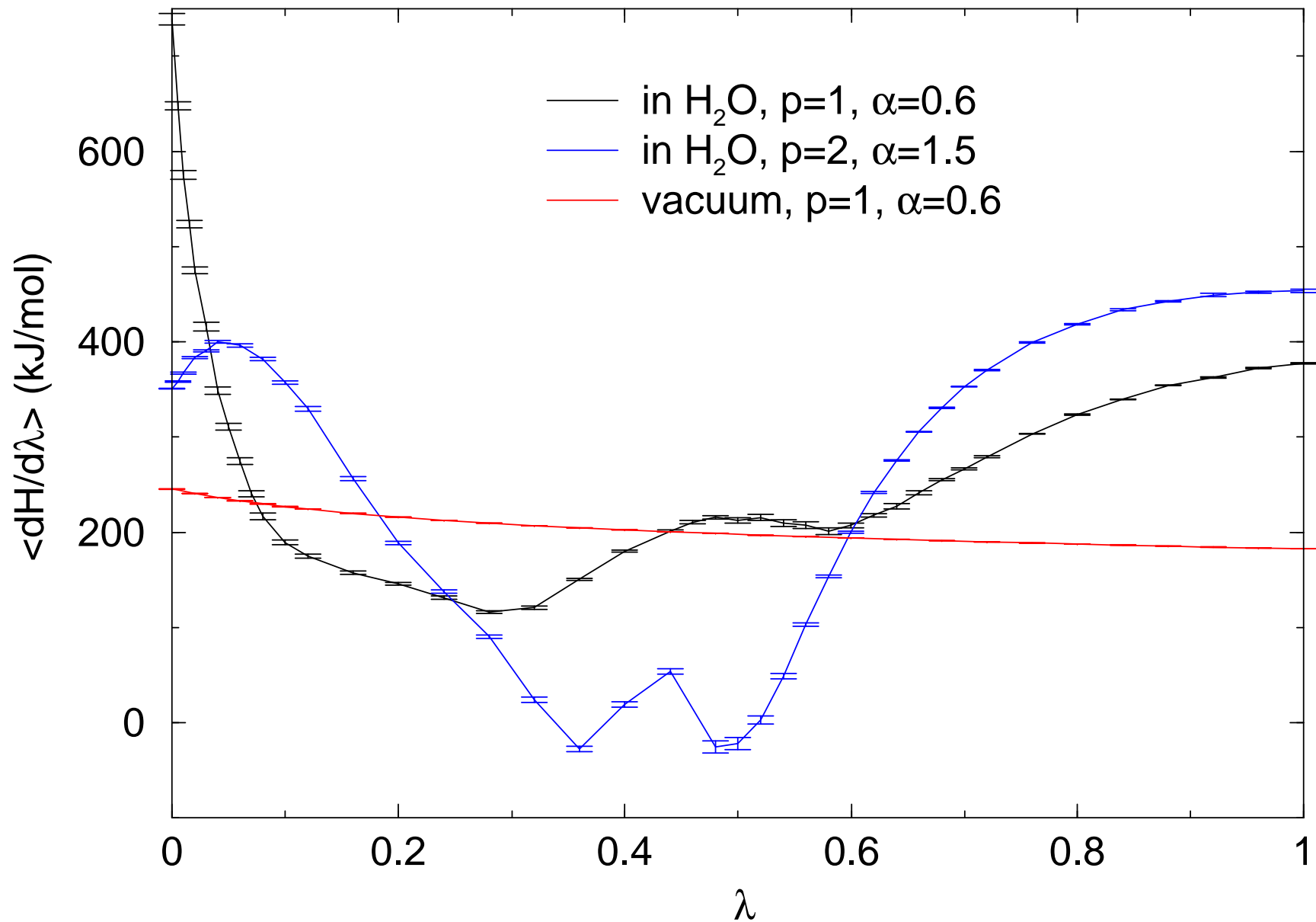
Turning off non-bonded interactions

In most cases it is more efficient to also turn off all non-bonded intramolecular interactions

(currently this is the only possibility in GROMACS)



Solvation of ethanol in water



Free energy topology

```
; Include forcefield parameters
#include "ffoplsaa.itp"
```

```
[ moleculetype ]
```

```
; Name          nrexcl
methanol        3
```

```
[ atoms ]
```

```
;nr  type  resnr  res  atom  cgnr  charge  mass  typeB  chargeB  massB
 1  opls_080  1  MET  CH3   1    0.266  15.035  opls_068  0  15.035
 2  opls_078  1  MET  O     1   -0.674  15.9994  opls_071  0  14.027
 3  opls_079  1  MET  H     1    0.408   1.008  opls_068  0  15.035
```

```
[ bonds ]
```

```
; ai  aj  funct  dA  kA  dB  kB
  1   2    1
  2   3    1
```

```
[ angles ]
```

```
; ai  aj  ak  funct  thetaA  kA  thetaB  kB
  1   2   3    1
```

Potential of mean force

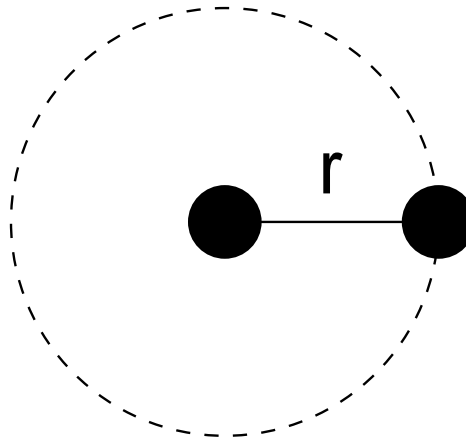
A potential of mean force (PMF) is the free energy along one or more degrees of freedom

The name comes from a common way to derive a PMF: by integrating the mean force working on a certain degree of freedom

One can also obtain a potential of mean force by Boltzmann inverting a pair correlation function $g(r)$:

$$PMF(r) = -k_B T \log(g(r)) + C$$

Entropic effects



The phase-space volume available to the system is: $4\pi r^2$

Thus the entropic distribution is:

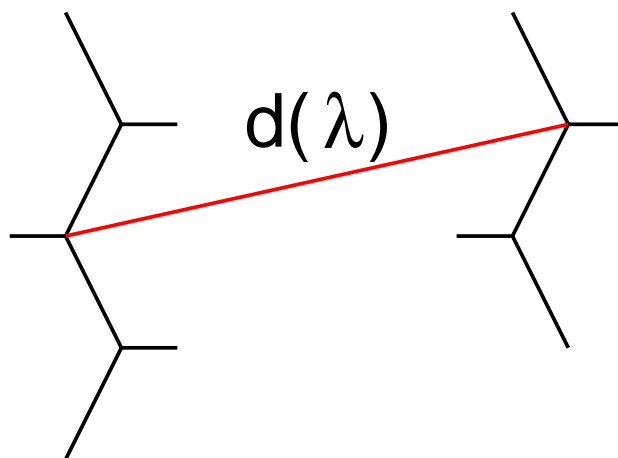
$$T\Delta S = k_B T \log(4\pi r^2) = 2k_B T \log(r) + C$$

This result can also be obtained by integrating the centrifugal force

$$F_c(r) = \frac{2k_B T}{r}$$

PMF with a coupling parameter

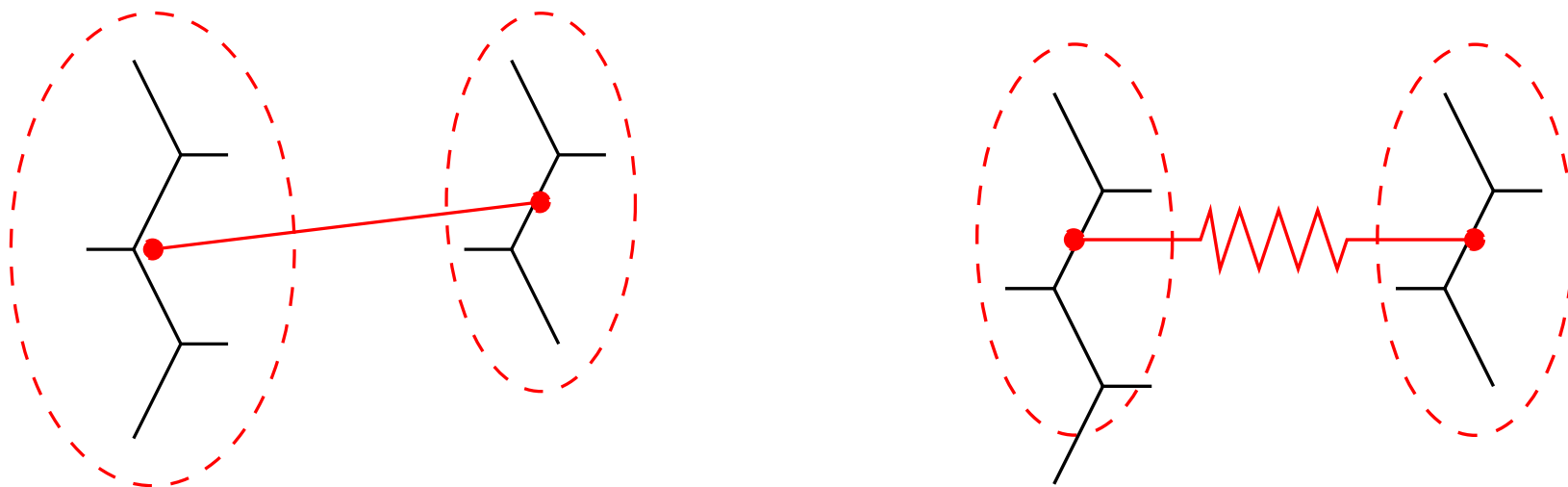
Since constraints are part of the Hamiltonian, they can also be coupled with lambda



The mean force is given by: $-\left\langle \frac{\partial H}{\partial \lambda} \right\rangle$

With this approach one can one make PMF for distances between atoms, not between centers of mass

PMF with the pull code

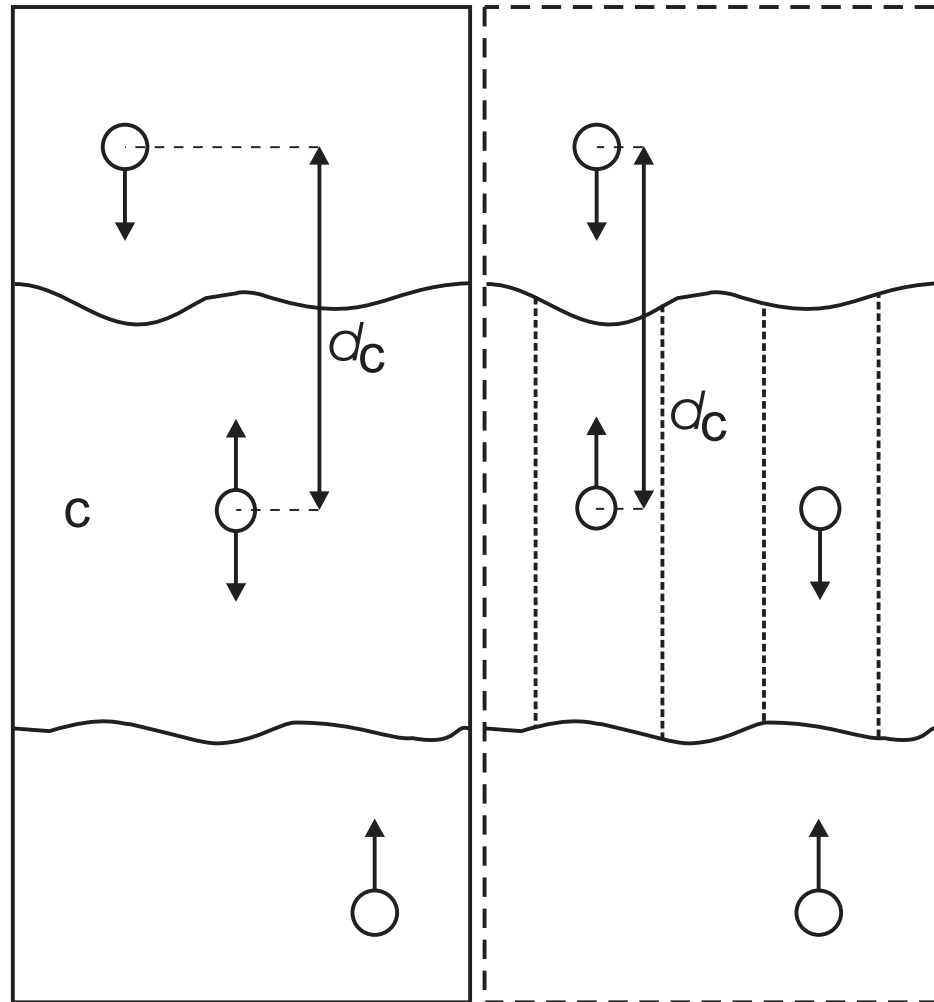


With a constraint: PMF between whole molecules

With a spring (umbrella sampling): PMF between anything
The mean force is the mean force working on the spring

Confusingly there are similar options *AFM* and *umbrella*

The pull code for membranes



The pull code in practice

The pull code is controlled by extra input files for `mdrun`

An index file to define the groups

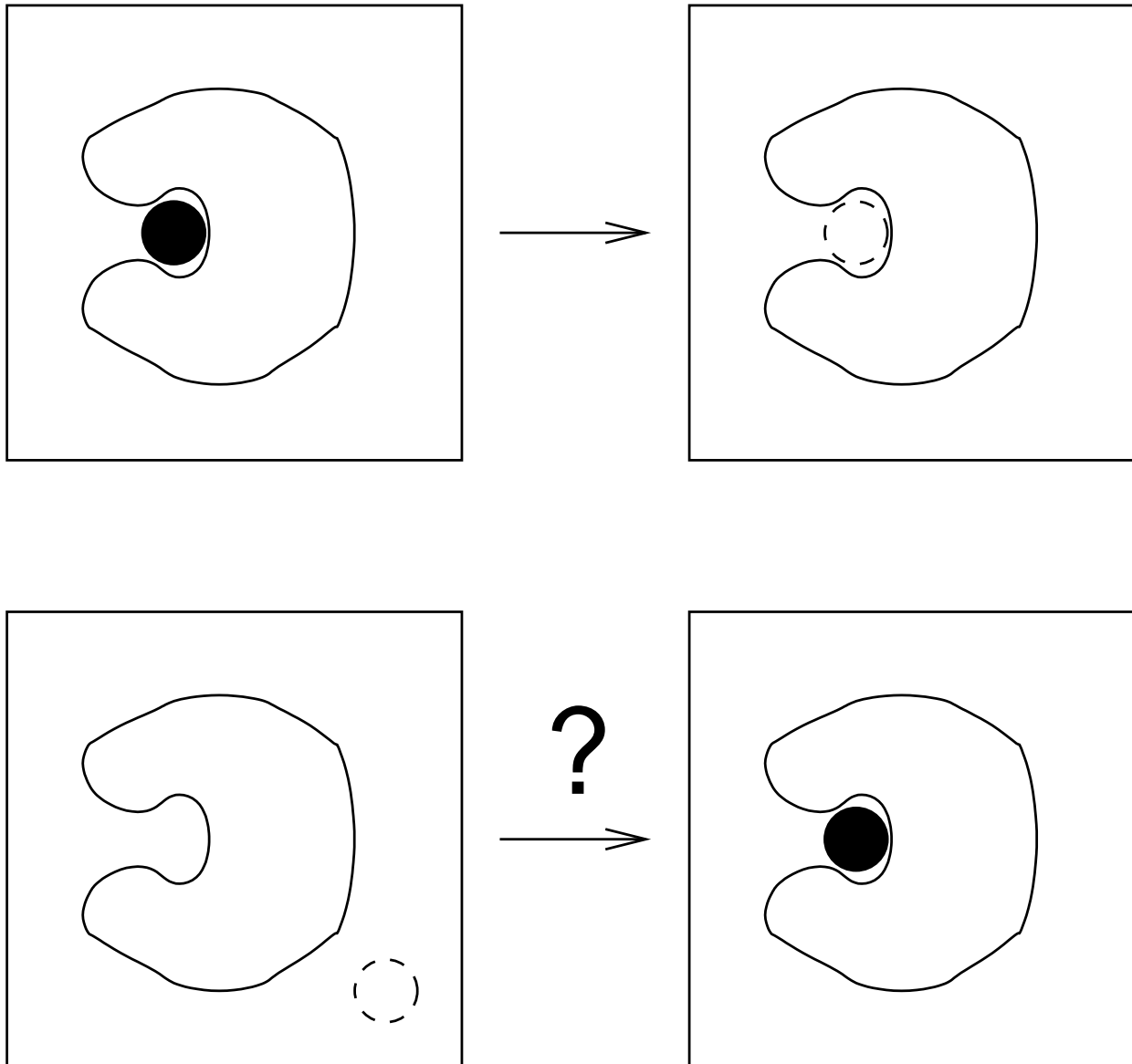
An `.ppa` file for the pull parameters:

Constraint pulling: constraint length, pull rate

“Spring” pulling: distance, pull rate, force constant

Output: `pull.pdo` with the pull forces

Reversibility



Reversibility check

The forward and backward path should give the same answer

So one can check:

$$\begin{aligned} \Delta G \text{ making molecule disappear} \\ = \\ -\Delta G \text{ make molecule appear} \end{aligned}$$

or

$$\begin{aligned} \Delta G \text{ pulling molecule out of a protein} \\ = \\ -\Delta G \text{ pushing molecule into a protein} \end{aligned}$$

Estimating errors

Block averaging

Estimating error is important for any quantitative analysis

A simple method that can always be applied is to split your data into m blocks and estimate the error by assuming the blocks are independent

Say we have n data points: $x(i\Delta t)$, $i = 0, \dots, n - 1$

We can estimate the error of the average from m block averages B_j :

$$E = \sqrt{\frac{1}{m(m-1)} \sum_{j=0}^{m-1} (B_j - \langle B \rangle)^2}$$

$$B_j = \frac{1}{\ell} \sum_{i=j\ell}^{(j+1)\ell-1} x(i\Delta t) \quad , \quad \ell = \frac{n}{m}$$

Block averaging

But the blocks will always be correlated

We can assume a double exponential correlation,
fast decay correlation time τ_1 , slow decay τ_2 :

$$\langle x(0) x(t) \rangle = a e^{-t/\tau_1} + (1 - a) e^{-t/\tau_2}$$

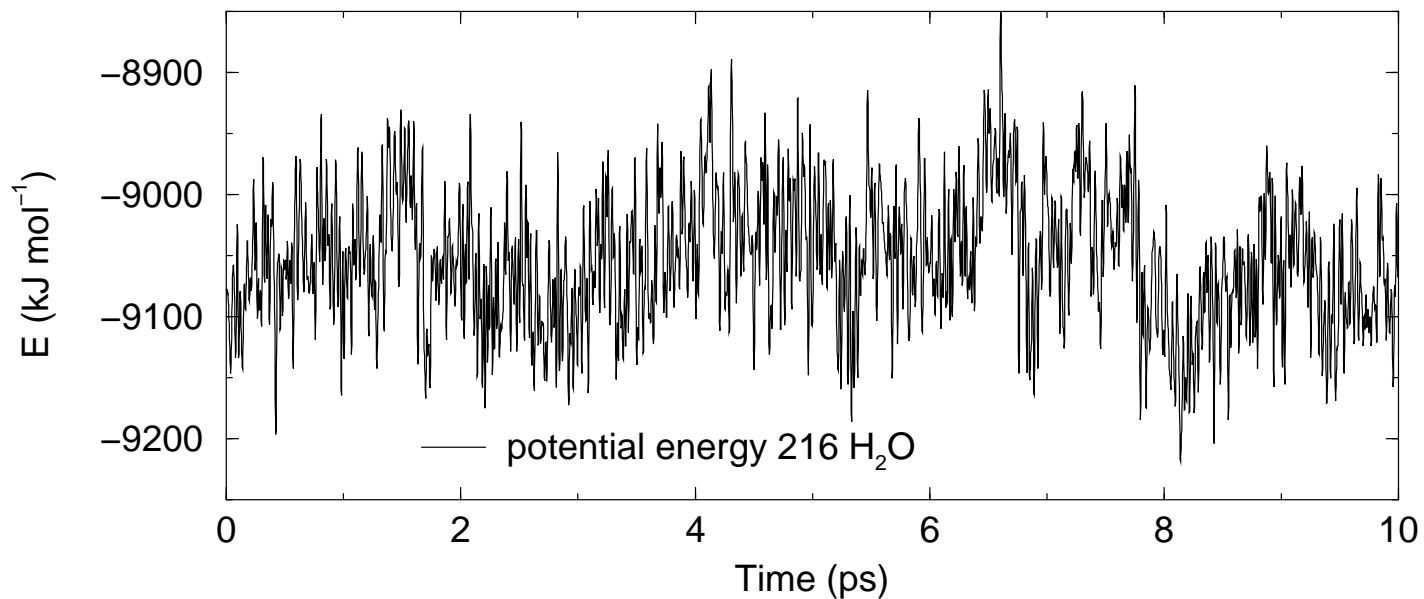
$$E^2(t) = \sigma^2 \frac{2}{n\Delta t} \left\{ a \tau_1 \left[(e^{-t/\tau_1} - 1) \frac{\tau_1}{t} \right] + (1 - a) \tau_2 \left[(e^{-t/\tau_2} - 1) \frac{\tau_2}{t} \right] \right\}$$

The error estimate for the average is then given by:

$$\lim_{t \rightarrow \infty} E(t) = \sigma \sqrt{\frac{2}{n\Delta t} \{ a \tau_1 + (1 - a) \tau_2 \}}$$

This analysis can be done with `g_analyze -ee`

Error estimate example



$$a = 0.72$$
$$\tau_1 = 0.016 \text{ ps}$$
$$\tau_2 = 0.50 \text{ ps}$$

