2. FREE ENERGY PERTURBATION (FEP)

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1 Free Energy Perturbation (FEP)

Free energy perturbation simulations can be used for theoretical prediction of binding free energies and along with it the elucidation and/or prediction of chemical reaction mechanisms. In this step-by-step tutorial you will be able to compute the relative binding free energy of camphane and camphor to an enzyme which catalyzes the addition of molecular oxygen to nonactivated hydrocarbons at physiological temperature. This oxydizing enzyme is called cytochrome P450 cam [1].

1.1 Theoretical background

The Free Energy Perturbation method is just that, a method, it is not, by any means, a theory. The method, in broad lines, allows a travel between states given a forced path. This makes sense since, by definition, the Free Energy of a system is a thermodynamic state function and therefore its value must be independent of the path taken. An approximation is made that a partition function for a perturbed system can be separated from the full partition function of the system as given by equation 3. The original expressions given by Zwanzing [2] to describe the idea of the Free Energy Perturbation (FEP) method as applied to hot gases are given below:

$$exp(-\beta \bar{A}_N) = Q_N = \frac{1}{N!} \int_{v} \dots \int_{v} \times exp(-\beta V_N) d_{\tau 1} \dots d_{\tau N}$$
(1)

$$V_N = V_N^{(0)} + V_N^{(1)} (2)$$

$$exp - \beta(\bar{A}_N - \bar{A}_N^{(0)}) \equiv exp(-\beta \bar{A}_N^{(1)})$$
 (3)

$$exp(-\beta \bar{A}_N^{(1)}) = \langle exp(-\beta V_N^{(1)}) \rangle_0 \tag{4}$$

Where $\beta=1/\kappa T$, κ is Boltzmann's constant, T is the temperature in Kelvin, Q_N is the configurational partition function of the system, \bar{A} is the configurational free energy, and V_N is the total potential energy of the system.

Notice that the paper is called "High Temperature Equation of State by a Perturbation Method. I. Nonpolar Gases". Why the title specifically states the high temperature case is because it deals with a gas where particles are not interacting with each other and so can be considered to be having fully elastic collisions.

A more modern notation to describe the perturbation method as applied to free energies is given by Pohorille, Jarzynski and Chipot[3] in the following expression:

$$exp(-\beta \Delta A) = \langle exp(-\beta \Delta U) \rangle \tag{5}$$

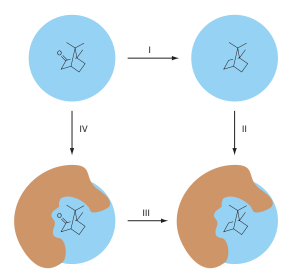


Figure 1: Thermodynamic cycle for ligand binding of camphene (CMA) and camphor (CAM) to the P450cam enzyme. The brown blob denotes the enzyme, and the blue background represents water.

Eq. 4 is derived from the statistical mechanics expression for the Helmholtz free energy, A, and the configurational integral, Z.

$$A = -kT \cdot \ln(Z) \tag{6}$$

$$Z = \int e^{-V/(kT)} d\Gamma \tag{7}$$

At constant pressure and volume the change in Helmholtz free energy equals that in Gibbs free energy. The free energy difference between the two different states A and B, that are represented with potential energies V_A and V_B respectively, can be rearranged into the eq. 4 as follows:

$$\Delta G_{A \to B} = -(kT \cdot \ln(Z_B) - kT \cdot \ln(Z_A))$$

$$= -kT \cdot \ln(Z_B/Z_A)$$

$$= -kT \cdot \ln(\left(\int e^{-V_B/(kT)} d\Gamma\right) / \left(\int e^{-V_A/(kT)} d\Gamma\right))$$

$$= -kT \cdot \ln(\left(\int e^{-V_B/(kT)} e^{-V_A/(kT)} e^{V_A/(kT)} d\Gamma\right) / \left(\int e^{-V_A/(kT)} d\Gamma\right))$$

$$= -kT \cdot \ln \langle e^{-\Delta V_{A \to B}/(kT)} \rangle_A$$
(8)

FEP is very accurate concerning small perturbations, when the difference between the potential V_A and V_B is smaller than 2 kT. The free energy of association of the ligands to a protein is calculated with FEP as a relative difference between the ligands. The absolute binding energy is defined as a free energy associated with moving the ligand in water to a solvated protein, paths II and IV in Figure 1. The relative binding energies are defined as a difference between two simulations where the ligand A is stepwise transformed to ligand B in water and in solvated protein, paths I and III in Figure 1.

1.2 FEP simulations using qdyn

Before starting the tutorial change the directory to protein or water in the FEP folder.

In this tutorial we are going to analyze the perturbation of camphor (CAM) to camphane (CMA) in water and in P450cam. P450 enzymes catalyze the hydroxylation of unactivated alkanes, and P450cam catalyzes specifically the hydroxylation of camphor. Although the system investigated here is P450cam the rules in this tutorial are general and can be applied to whatever system of interest.

A FEP simulation is carried out in **Q** with a name.fep file where the transformation is defined in detail. Every name.fep is divided in a number of sections where we define atoms, charges, bonds, angles etc that are changing during the FEP simulation (Figure 2). Note that a section in **Q** follows the syntax [...].

- Open the cma_cam.fep, in the protein or water folder, that is used for the CAM to CMA perturbation. Try to locate the different sections that are used in the FEP file. Can you understand how CMA is transformed to CAM?
- What effect do they have on the simulation? (HINT: For the detailed explanation of the different sections consult Appendix A!)

```
[atoms]
...
[FEP]
...
[change_charges]
...
[atom_types]
...
.
```

Figure 2: FEP file format.

Having a FEP file the perturbation is defined in Q by adding a specifier

```
fep name.fep
```

in the input files for **qdyn**. Additionally it is necessary to add the section [lambdas] where the lambda values are defined, λ_1 and λ_2 . The lambda values are used to transform the potential V_A to V_B in small steps to improve the convergence. The energies are sampled on the potential V,

$$V = \lambda_1 V_A + \lambda_2 V_B = \lambda_1 V_A + (1 - \lambda_1) V_B \tag{9}$$

where λ_1 varies between 0 and 1. How many lambda steps are used in the simulation depends on difference between V_A and V_B and the type of perturbation.

- Open an input file name.inp. In what section is the fep file read?
- Find the [lambdas] section. How many lambda steps are used in the perturbation and how big is each step? (HINT: Look in several cma_camN.inp files, where N=0..30.)

- How many steps are used in the simulation in each file?
- Also note the specifiers

energy name.en energy 25

that are used for saving the energies every 25th step to the file name.en.

1.3 Analysis of FEP simulations using qfep

All steps in this tutorial take approximately 2-3 hours running in parallel in eight processor cores in a 2.3 GHz processor. **qfep** is the analysis program which allows you to compute the free energy differences between states using energies which have been stored in the binary files with ".en" extension, that is, files such as cma_cam30.en. To run qfep you must issue the following command for both the simulation in plain water and in the protein:

The qfep.inp file contains all necessary instructions issued to **qfep** and the qfep.out contains the full output.

- Open qfep.inp and qfep.out. Try to understand the different commands. What are they specifying? (HINT: Check out Appendix B!)
- The free energy difference for the perturbation from CMA to CAM can be found at the end of table summarized in Part 1. What is the difference between water and protein?

Water simulation: $\Delta G_{A \to B}^w = (\text{kcal/mol})$

Protein simulation: $\Delta G^p_{A \to B} = (\text{kcal/mol})$

The free energy is calculated as an average of forward and backward calculations at every lambda point. The forward and backward free energy values are also summarized in Part 1 of the qfep.out file in columns sum(dGf) and sum(dGr). The theoretical error of the FEP simulation is half the value of the difference between forward and backward calculations.

-What is the theoretical error of FEP for water and protein simulations?

 $\mbox{Water simulation: } \Delta G_{A \rightarrow B}^{w,error} = \mbox{(kcal/mol)}$

Protein simulation: $\Delta G_{A \to B}^{p,error} =$ (kcal/mol)

The relative free energy of binding is determined as the difference between the free energies for the protein and water simulations.

Relative binding free energy:

Calculated: $\Delta\Delta G_{bind,rel}^{calc} = \Delta G_{A\to B}^{p} - \Delta G_{A\to B}^{w} =$ (kcal/mol)

Experimental: $\Delta\Delta G^{exp}_{bind.rel} = \Delta G^p_{A \to B} - \Delta G^w_{A \to B} =$ -2.0 (kcal/mol)

Alternatively the free energy profiles can be plotted and investigated graphically. The program <code>gnuplot</code> is used for plotting different graphs.

- Open gnuplot by typing gnuplot in the shell window and then write load 'fep.part1.pgp'. This script plots the free energy as a function of lambda, as summarized in the table in Part 1 of qfep.out.
 - What is the shape of the curve? What is the spacing between the points? Does it look ok?

Molecular dynamics simulations also gives us an opportunity to investigate the structures as they are propagated through time.

- Type **pymol** and open cma_cam.pse which loads cma_cam0.pdb, cma_cam10.pdb and cma_cam30.pdb files in **pymol**. The name.pdb files represent structures at the lambda state (1,0), (0.63,0.37) and (0,1) respectively.
 - What is the difference between the different ligand structures?

- What are the interactions between the ligands and the surrounding. (HINT: Check out Tyr87 residue!)

Here ends the FEP part of the lab course!

A FEP file format

Table 1: FEP file format

[atoms]: Define Q-atoms.

column	description
1	Q-atom number (counting from 1 up).
2	Topology atom number.

[PBC]: For periodic boundary conditions.

keyword	value	comment
switching-	Topology atom number.	Required with periodic boundary condi-
_atom		tions.

[FEP]: General perturbation information.

keyword	value	comment
states	Number of FEP/EVB states.	Optional, default 1.
offset	Topology atom number.	Optional, default 0. This number is added
		to all topology atom numbers given in the
		FEP file.
offset_residue	Residue/fragment number.	Optional. Set offset to the topology num-
		ber of the first atom in the given residue
		minus one.
offset_name	Residue/fragment name.	Optional. Set offset to the topology num-
		ber of the first atom in the first residue with
		the given name minus one.
qq_use-	This is a special feature for studying $e.g.$ elec-	Optional, default off.
_library-	trostatic linear response. Set to 'on' to use the	
_charges	library charges from the topology for intra-Q-	
	atom interactions, i. e. change only Q-atom-	
	surrounding electrostatic interactions.	
softcore-	Set to 'on' if the values entered in the [softcore]	Optional, default off.
_use_max-	section are the desired maximum potentials	
_potential	(kcal/mol) at $r=0$. Qdyn will then calculate	
	pairwise α_{ij} to be used in equation 10. 'off'	
	means the values are to be used directly in	
	equation 10.	

[change_charges]: Redefine charges of Q-atoms.

column	description
1	Q-atom number (referring to numbering in atoms section).
2	Charge (e) in state 1, state 2,

[atom_types]: Define new atom types for Q-atoms: Standard LJ parameters and parameters for the exponential repulsion potential $U_{soft}=C_i\cdot C_j\epsilon^{-a_i\cdot a_j\cdot r_{i,j}}$.

1	Name (max 8 characters).
2	Lennard-Jones A parameter $(kcal^{\frac{1}{2}} \cdot mol^{-\frac{1}{2}} \cdot \mathring{A}^{-6})$ for geometric combination or R* $(kcal \cdot mol^{-1} \cdot \mathring{A}^{-12})$ for arithmetic combination rule.
3	LJ B parameter (kcal $\frac{1}{2}$ ·mol $-\frac{1}{2}$ ·Å -3) or ϵ (kcal·mol -1 ·Å -6).
4	Soft repulsion force constant C_i (kcal $^{\frac{1}{2}}$ ·mol $^{-\frac{1}{2}}$) in U_{soft} .
5	Soft repulsion distance dependence parameter \mathbf{a}_i ($\mathbf{\mathring{A}}^{-\frac{1}{2}}$) in U_{soft} .
6	Lennard-Jones A parameter $(kcal^{\frac{1}{2}} \cdot mol^{-\frac{1}{2}} \cdot \mathring{A}^{-6})$ or R^* $(kcal \cdot mol^{-1} \cdot \mathring{A}^{-12})$ for 1-4 interactions.
7	LJ B parameter (kcal $\frac{1}{2}$ ·mol $-\frac{1}{2}$ ·Å -3) or e (kcal·mol -1 ·Å -6) for 1-4 interactions.
8	Atomic mass (u).

[change_atoms]: Assign Q-atom types to Q-atoms.

[onango_atome	[onango_atomo]. Addigit a atom types to a atomo.	
1	Q-atom number.	
2	Q-atom type name in state 1, state 2,	

[soft_pairs]: Define pairs which use soft repulsion.

1	Q-atom number of first atom in pair.
2	Q-atom number of second atom in pair.

[excluded_pairs]: Define pairs to exclude from non-bonded interactions. Note: also non-Q-atoms can be excluded.

1	Topology atom number of first atom in pair.
2	Topology atom number of second atom in pair.
3	Exclusion effective (1) or not (0) in state 1, state 2,

[el_scale]: Define q-atom pairs for scaling of the electrostatic interaction. Can be useful e.g. when highly charged intermediates appear in FEP/EVB. The scale factor applies to all states. Note: only Q-atom pairs can be scaled.

1	q-atom number of first atom in pair
2	q-atom number of second atom in pair
3	electrostatic scale factor (01)

[softcore]: Define q-atom softcore potentials. The meaning of these entries depends on the value of softcore_use_max_potential.

1		q-atom number
2		Desired potential at $r=0$ for all of this q-atom's vdW interactions in state 1, state 2, or
		the actual α value used in equation 10. An α of 200 yields vdW potentials at $r=0$ of 10-50
		kcal/mol for heavy atom - heavy atom interactions. Set to 0 if softcore is not desired for this
		q-atom.

[monitor_groups]: Define atom groups whose non-bonded interactions are to be monitored (printed in the log file).

		T 1
1 1		Topology atom number of first and following atoms in group.
		I TODOTOUV ATOTTI HUTTIDEL OF III SEATU TOTOWITU ATOTTIS III UTOUD.
1	-	1

[monitor_group_pairs]: Define pairs of monitor_groups whose total non-bonded interactions should be calculated.

1	First monitor_group number.
2	Second monitor_group number.

[bond_types]: Define Q-bond types using Morse or harmonic potentials,

$$E_{Morse} = D_e \left(1 - e^{-\alpha(r-r_0)} \right)^2 E_{Harmonic} = \frac{1}{2} k_b (r - r_0)^2.$$

Morse and harmonic potentials can be mixed (but each bond type is either kind). Entries with four values are Morse potentials and entries with three values are harmonic.

	Morse potential	Harmonic potential
1	Q-bond type number (s	starting with 1).
2	Morse potential dissociation energy, D_e (kcal·mol ⁻¹).	Harmonic force constant k_b (kcal·mol ⁻¹ ·Å ⁻²).
3	Exponential co-efficient α in Morse potential (\mathring{A}^{-2}) .	Equilibrium bond length r_0 in harmonic potential ($\mathring{\mathbf{A}}$).
4	Equilibrium bond length r_0 in Morse potential (Å).	

[change_bonds]: Assign Q-bond types. Note: shake constraints for the redefined bonds are removed. The order in which atoms are given is not important.

1	Topology atom number of first atom in bond.
2	Topology atom number of second atom in bond.
3	Q-bond type number (referring to numbering in bond_types section) or 0 to disable bond in state 1, state 2,

[angle_types]: Define Q-angle types.

	. 5 _ 71	<u> </u>
	1	Q-angle type number (starting with 1).
ĺ	2	Harmonic force constant $(kcal \cdot mol^{-1} \cdot rad^{-2})$.
	3	Equilibrium angle (°).

[change_angles]: Assign Q-angle types.

1	Topology atom number of first atom in angle.
2	Topology atom number of middle atom in angle.
3	Topology atom number of third atom in angle.
4	Q-angle type number (referring to numbering in angle_types section) or 0 to disable angle
	in state 1, state 2,

[torsion_types]: Define Q-torsion types.

1	Q-torsion type number (starting with 1).
2	Force constant = $\frac{1}{2}$ ·barrier height (kcal·mol ⁻¹).
3	Periodicity (number of maxima per turn).
4	Phase shift (°).

[change_torsions]: Assign Q-torsion types. Note: The order of atoms (1, 2, 3, 4 or 4, 3, 2, 1) is not important.

1	Topology atom number of first atom in torsion.
2	Topology atom number of second atom in torsion.
3	Topology atom number of third atom in torsion.
4	Topology atom number of fourth atom in torsion.
5	Q-torsion type number (referring to numbering in torsion_types section) or 0 to disable
	torsion in state 1, state 2,

[improper_types]: Define Q-improper types.

-	1	Q-improper type number (starting with 1).
2	2	Harmonic force constant (kcal·mol $^{-1}$ ·rad $^{-2}$).
3	3	Equilibrium angle (°).

[change_impropers]: Assign Q-improper types. Note: The order of atoms (1, 2, 3, 4 or 4, 3, 2, 1) is not important.

1	Topology atom number of first atom in improper.
2	Topology atom number of second atom in improper.
3	Topology atom number of third atom in improper.
4	Topology atom number of fourth atom in improper.
5	Q-improper type number (referring to numbering in improper_types section) or 0 to disable
	improper in state 1, state 2,

[angle_couplings]: Couple Q-angles to Q-bonds, *i.e.* scale angle energy by the ratio of the actual value of the Morse bond energy to the dissociation energy.

1	Q-angle number (line number within change_angles section).
2	Q-bond number (line number within change_bonds section).

[torsion_couplings]: Couple Q-torsions to Q-bonds.

1	Q-torsion number (line number within change_torsions section).
2	Q-bond number (line number within change_bonds section).

[improper_couplings]: Couple Q-impropers to Q-bonds.

1	Q-improper number (line number within change_impropers section).
2	Q-bond number (line number within change_bonds section).

[shake_constraints]: Define extra shake constraints. The effective constraint distance will be the sum of the distances given for each state, weighted by their I values. Note: constraints defined here do not override constraints imposed by setting the shake flag to *on* in the Qdyn input file. To remove a constraint the bond must be redefined as a Q-bond. The order in which atoms are given is not important.

1	Topology atom number of first atom.
2	Topology atom number of second atom.
3	Constraint distance (Å) in state 1, state 2,

[off-diagonals]: Define off-diagonal elements of the Hamiltonian, represented by $H_{i,j}=A_{i,j}\cdot\epsilon^{-\mu_{i,j}\cdot r_{k,l}}$ where i and j are states and k and l are Q-atoms.

1	State i.
2	State j.
3	Q-atom k.
4	Q-atom I.
5	$A_{i,j}$ (kcal·mol ⁻¹).
6	$\mu_{i,j}$ (Å $^{-1}$).

Softcore equation:

$$V_{vdW}(r) = \frac{A_{ij}}{(r^6 + \alpha_{ij})^2} - \frac{B_{ij}}{r^6 + \alpha_{ij}} \qquad \text{or} \qquad V_{vdW}(r) = \epsilon \cdot \left(\frac{R_{ij}^{* \ 12}}{(r^6 + \alpha_{ij})^2} - 2 \cdot \frac{R_{ij}^{* \ 6}}{r^6 + \alpha_{ij}}\right)$$
(10)

B qfep input summary

Some of the commands are affecting only analysis of the empirical valance bond (EVB) simulations, while others are common for both FEP and EVB. The ones affecting FEP are marked in bold text and are of concern in FEP simulations.

HINT:

-> specifies the input commands # denotes the written output.

```
->Number of energy files:
# Number of files = 31
-> No. of states, no. of predefined off-diag elements:
# Number of states = 2
# Number of off-diagonal elements = 0
-> Give kT & no, of pts to skip:
\# kT = 0.596
# Number of data points to skip = 80
-> Give number of gap-bins:
# Number of gap-bins = 40
-> Give minimum # pts/bin:
# Minimum number of points per bin= 10
-> Give alpha for state 2:
# Alpha for state 2 = 0.00
-> Number of off-diagonal elements:
# Number of off-diagonal elements = 0
-> linear combination of states defining reaction coord:
# Linear combination co-efficients= 1.00 0.00
name1.en
name2.en
nameN.en
```

The # Number of files is the total number of files used for the FEP simulation, # Number of data points to skip is the number of points that are discarded as the equilibration at each lambda step and # kT specifies the temperature of the simulation. The energy files name.en are read last in **gfep**.

References

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