

Poly-Proline with AMBER 11.

V. Babin and C. Sagui

April 27, 2010

Abstract

This document describes a couple of ways to assess the conformational equilibria of a short polyproline peptide using `sander.MPI` program from the AMBER package version 11. In particular, so called *steered molecular dynamics* is illustrated along with the famous *replica-exchange*[1] protocol. The familiarity with the subject is assumed as the intent of this document is to expose the technicalities rather than describe the basics.

1 Introduction.

When preparing to write this note, I faced the following unavoidable choice: (1) use a simple and hence relatively boring example that is easily runnable within an hour or (2) consider a larger system that takes longer to run, yet is marginally more interesting. Option (2) won this time, and hence this text describes two ways of simulating the $\text{Ace}-(\text{Pro})_5-\text{NHe}$ peptide in AMBER 11. All the files and scripts needed to reproduce the runs described in this note should be available alongside with this document in the `polyproline-tutorial-files.tar.bz2` archive (Warning: ≈ 60 Mb).

2 Preliminaries.

We make the `prmtop` and `inpcrd` files using the `teLeap` program with the input shown in Fig.1. We are going to use GB/SA solvation model from Refs.[2, 3, 4] (`igb = 2` and `gbsa = 1`) and therefore line 5 instructs the `teLeap` program to use appropriate radii. The rest should be self-explanatory.

It is instructive to try regular molecular dynamics to see the lack of ergodicity at room temperature (assumed to be 300K). To this end, one can use the `mdin` file shown in Fig.2. Apart from the usual `&cntrl` namelist the file includes the `ncsu_abmd` section that defines a *collective variable* of type of `COS_OF_DIHEDRAL` (lines 17–25) and instructs `sander` to save the values of the variable every 500 steps (1 ps) to the `monitor.txt` file (lines 12, 14, and 15). The collective variable is given by the sum of cosines of the dihedral angles along the peptide bonds connecting successive proline residues (see comments in Fig.2, lines 20–24). For all the bonds in *cis* conformation, the variable would be equal to five, while for the all *trans* configuration, it would be equal minus five. Once

```
1 source leaprc.ff99SB
2
3 m = sequence {ACE PRO PRO PRO PRO PRO NHE}
4
5 set default PBradii mbondi2
6 saveamberparm m prmtop inpcrd
7 savepdb m inpcrd.pdb
8
9 quit
```

Figure 1: Input file for the LEaP program used to generate the prmtop and inpcrd files for the Ace – (Pro)₅ – NHe peptide.

the simulation is complete, examination of the `monitor.txt` file shows that no *cis/trans* transitions happened over one nanosecond. It is instructive to run it longer and try higher temperatures as well. Summarizing, the regular molecular dynamics is not ergodic in this case: the trajectory does not visit all equilibrium configurations with Boltzmann probability; it gets stuck in the vicinity of the initial configuration because of the high (free) energy barriers separating different (free) energy minima. The aim of this tutorial is to show two possible routes to study the equilibrium ensemble in spite of presence of these barriers. In the next section we are going to consider so-called *steered molecular dynamics* that can be used to compute the free energy differences between different conformations. Afterwards we are going to go through a more demanding (yet more powerful as well) *replica-exchange*[1] technique.

The files described above can be found in the `polyproline-tutorial-files/2.preliminaries` directory.

3 Steered molecular dynamics.

The so-called *steered molecular dynamics* amounts to a non-stationary harmonic restraint applied to a (set of) collective variable(s). The method is extensively discussed in the literature and we therefore skip any details here directing the reader to the excellent Refs.[5, 6, 7] and references therein. A short summary for the lazy: one can “pull” the system between different states, measure the non-equilibrium work performed, and then infer corresponding *equilibrium* free energy differences.

Several subsystems in `sander` implement the “pulling” functionality and only one of them (`ncsu_smd`) is exposed in this writeup. Curious reader is directed to the user manual to find out more on these and related matters.

We proceed by pulling on the `COS_OF_DIHEDRAL` variable described above changing its value between -5 and 5 (that is, between the all-*trans* and all-*cis* conformations). To get a robust Δf we perform one hundred of forward pullings (going from -5 to +5) and one more hundred of reverse pullings (going from +5 to -5). Initial coordinates used for each of the `ncsu_smd` runs were equilibrated

```

1 **** this line is required ****
2 &cntrl
3   ntwx = 0, ntp = 5000, ntwr = 50000,
4   ntt = 3, temp0 = 300.0, gamma_ln = 1.0,
5   igb = 2, gbsa = 1, dielc = 1.0, cut = 18.0,
6   ntb = 0, ntc = 2, ntf = 2, tol = 0.000001,
7   nstlim = 500000, dt = 0.002, ntp = 0, ibelly = 0,
8   ntr = 0, imin = 0, irest = 0, ntx = 1, ig = 27606
9 /
10
11 ncsu_abmd
12   mode = ANALYSIS
13
14   monitor_file = 'monitor.txt'
15   monitor_freq = 500 ! 1 ps
16
17   variable
18     type = COS_OF_DIHEDRAL
19
20     i = ( 2, 5, 7, 17, ! :1@CH3 == :1@C == :2@N == :2@CA
21           17, 19, 21, 31, ! :2@CA == :2@C == :3@N == :3@CA
22           31, 33, 35, 45, ! :3@CA == :3@C == :4@N == :4@CA
23           45, 47, 49, 59, ! :4@CA == :4@C == :5@N == :5@CA
24           59, 61, 63, 73) ! :5@CA == :5@C == :6@N == :6@CA
25   end variable
26 end ncsu_abmd

```

Figure 2: Input file for the `sander` program suitable for doing regular molecular dynamics at room temperature.

in the corresponding state (all-*trans* for the forward pullings, and all-*cis* for the reverse pullings). Relevant section of `mdin` file is shown in Fig.3 and should be self-explanatory. The progress of the pulling is reported to the `'work.txt'` file every 10 ps. The total work is reported at the end of this file once the simulation is complete. The files for all two hundred pulls can be found in `polyproline-tutorial-files/3.smd` directory and we encourage the reader to examine (or re-run) them. There is also a perl script that can be used to compute the free energy difference.

Numerical values of the non-equilibrium work for the forward (W^F) and reverse (W^R) pullings can be used to compute the Δf estimate using Crooks[5] relations (the equation for the estimator coincides with the earlier Bennet's acceptance ratio formula[8]) from the following equation:

$$\sum_{i=1}^{n_F} \frac{1}{1 + n_F/n_R \exp(W_i^F - \Delta f)} - \sum_{i=1}^{n_R} \frac{1}{1 + n_R/n_F \exp(W_i^R + \Delta f)} = 0, \quad (1)$$

with $n_{F,R}$ denoting the numbers of forward and reverse simulations, respectively. We got 4.58 kcal/mol for the Δf . It is instructive to compare this estimator with the ones computed from the

Jarzynski[9] identity applied to the only forward (10.65 kcal/mol) or only reverse (-1.49 kcal/mol) processes.

```

1  **** this line is required ****
2  &cntrl
3      ...
4  /
5
6  ncsu_smd
7      output_file = 'output/00'
8      output_freq = 5000 ! 10 ps
9
10     variable
11         type = COS_OF_DIHEDRAL
12
13         i = ( 2, 5, 7, 17, ! :1@CH3 == :1@C == :2@N == :2@CA
14              17, 19, 21, 31, ! :2@CA == :2@C == :3@N == :3@CA
15              31, 33, 35, 45, ! :3@CA == :3@C == :4@N == :4@CA
16              45, 47, 49, 59, ! :4@CA == :4@C == :5@N == :5@CA
17              59, 61, 63, 73) ! :5@CA == :5@C == :6@N == :6@CA
18
19         path = (-5.0, 5.0) harm = (100.0)
20     end variable
21 end ncsu_smd

```

Figure 3: Input file for steered molecular dynamics going from all *trans* to all *cis* configuration of the Ace – (Pro)₅ – NHe peptide (the &cntrl namelist is not shown).

4 (Hamiltonian) replica exchange.

The *replica-exchange* method is a powerful formulation first proposed by Geyer in 1991 in conference proceedings[1] and subsequently rediscovered in a different area of science[10]. The basic idea of the technique is to exchange configurations between several simulations in a way that preserves detailed balance and hence makes every simulation to sample from its target distribution. The barriers that are high in one replica (referred to as “cold” replica with respect to *the* barriers) may happen to be significantly lower in another one (“hot” replica) so that the barriers crossing is greatly enhanced.

The success of the method depends crucially on the choice of the “hot” replica(s). Here we are going to use the *potentials of mean force* (PMF) to bias the dynamics in the “hot” replicas. The advantage of this approach over the more conventional *parallel tempering* is that the temperature does not change across the replicas hence rendering this kind of setup more suitable for explicit solvent simulations. We (shamelessly) refer curious readers to our preprint <http://arxiv.org/abs/0911.5132v2> for a more comprehensive presentation.

The idea is very simple (and by no means ours): assuming that a “slow mode” can be described by a collective variable $\sigma(\mathbf{r})$, and that the *potential of mean force* (also referred to as *Landau free energy*) associated with that collective variable is known, one can construct a “hot” replica by biasing the original dynamics by the negative of the PMF thus rendering the states with different values of $\sigma(\mathbf{r})$ equally probable. In practice the PMF is typically unknown, yet fortunately, a biasing potential that is within a few $k_B T$ of the true PMF, will do the trick. The latter can be computed in a variety of ways. In this writeup we are going to use the so-called ABMD[11] (adaptively biased molecular dynamics), which is yet another variation of the earlier local elevation[12] method (LEM), grown up from our efforts to make the so-called metadynamics[13, 14] method (another LEM variation) more practical.

For the $\text{Ace} - (\text{Pro})_5 - \text{NHe}$ peptide, one readily identifies the *cis/trans* transitions of the peptide bonds as the “slow modes”. There are five of them, and hence our aim is to setup a replica-exchange scheme with five “hot” replicas biased by the PMFs associated with the corresponding dihedral angles, five more replicas biased by the scaled down PMFs (to serve as proxies for more efficient random walk between the replicas), and one *not biased* replica, from which we are going to “read” the equilibrium samples. We compute the (approximate) PMFs using ABMD. For the very first, rough approximation (initial condition for more demanding replica-exchange runs), we assume that the PMF for the $:1\text{C}\text{H}3 == :1\text{C} == :2\text{N} == :2\text{C}\text{A}$ dihedral of the $\text{Ace} - (\text{Pro})_5 - \text{NHe}$ is close to the same PMF computed for $\text{Ace} - \text{Pro} - \text{NHe}$, and that the PMFs for the dihedrals between Pro residues in the $\text{Ace} - (\text{Pro})_5 - \text{NHe}$ are close to the one computed for $\text{Ace} - (\text{Pro})_2 - \text{NHe}$ dimer.

We begin with the monomer and proceed by doing two-stage “flooding”: starting with more aggressive (less equilibrium) and following it up with a smoother (closer to the equilibrium). Please find the corresponding files in the `4.hremd/1.preliminaries/0.monomer` subdirectory of the supporting files archive. The “flooding” here means naïve, linear grows of the biasing potential aimed to compensate for the true PMF. We refer the reader to our preprint for the details. Finally, we check the “quality” of the PMF by doing short *equilibrium* biased run (the files are in `0.monomer/3.umbrella`). It turns out that the angle indeed explored all possible values in this short run. This implies that the biasing potential is within few $k_B T$ of the true PMF, as intended. If it turned out to be not the case, we would go with slower and slower flooding as long as needed.

In the next step we compute PMF for the $:2\text{C}\text{A} == :2\text{C} == :3\text{N} == :3\text{C}\text{A}$ dihedral of $\text{Ace} - (\text{Pro})_2 - \text{NHe}$ dimer following the same routine (the files in `4.hremd/1.preliminaries/0.dimer`).

In order to equilibrate initial replica-exchange setup and to check whether those approximate PMFs computed using smaller molecules do approximate the true ones with the accuracy not worse than a few $k_B T$, we ran a short replica-exchange simulation comprising ten replicas (five fully biased and five “proxies”). These files are located in `4.hremd/2.hremd.eq` and we encourage the reader to examine them. We utilize the familiar `multisander` philosophy, yet use another (not mainstream) code path which is activated by the `ncsu_bbmd` section in the `mdin` files. Everything there should be self-explanatory (hit me by an e-mail, if it is not). This short run has revealed that the initial PMF do not approximate the true ones very well, as the values of the dihedrals are distributed very non-uniformly. To correct for that, we did a “flooding” run (`4.hremd/3.hremd.flooding`) with stationary “proxy” biasing potentials. We then checked the (presumably) more accurate PMFs by

doing a short equilibrium run (`4.hremd/4.hremd.umbrella`) and proceeded to the production phase (in case we were unhappy with the PMFs quality, we would do more of slow “flooding”).

The “production” run is in `4.hremd/5.hremd.production`. It comprises eleven replicas (five “hot”, five “proxies” and one “cold”) each ran for 50 ns. We try to exchange five random pairs of replicas every 91 step. The exchange statistics is reported in the `exchange_log_file` and turned out to be quite well.

Upon completion of the “production” run we got: (1) the monitor files from all eleven replicas: ten holding biased samples of the dihedral angles, and eleventh with the equilibrium samples of the `COS_OF_DIHEDRAL` collective variable; (2) unbiased configurations saved as the trajectory from the eleventh replica. We are going to analyze the trajectory, noting in passing that the biased samples can be used to improve the precision of the PMFs associated with the dihedral angles. We used `ptraj` to extract the values of the slow angles from the trajectory. It can be seen that the angles are indeed very sharply concentrated near the *cis* and *trans* values and hence they can be unambiguously classified as being either *cis* or *trans*. It is thus natural to use five-letter string consisting of T (for *trans*) and C (for *cis*) to describe each trajectory frame. A simple perl script (`omegas.pl` file in the `scripts` subdirectory of the supplementary archive) can be used to compare the frequencies of different conformations. We got that the TTTTT (all *trans*) configuration is indeed the most probable one (global minimum) with the probability of 33%. It is also possible to compute the free energy difference between the TTTTT and CCCCC states (that would correspond to the `COS_OF_DIHEDRAL` values of -5 and +5 respectively): we got 4.42 kcal/mol, which compares very favorably with the *steered molecular dynamics* described above.

References

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