One-Dimensional Weighted-Ensemble Brownian Dynamics Simulation in Perl

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Introduction

- Pharmaceuticals that cure illness, treat disease, and improve quality of life are the byproducts of our greater understanding of how the human body functions.

- In the last century, our knowledge of human anatomy and physiology has penetrated beyond the level of organ systems, organs, differentiated tissue cells, organelles, and nucleic acids to the very molecules of functionality themselves, proteins.

- It is through our better understanding of proteins the pharmaceutical companies have been able to eliminate or facilitate specific reactions in the body by targeting the proteins that act as their catalysts.
Background

- It is necessary to know a protein’s active state to develop drugs that will act as powerful inhibitors of protein function
  - It is often difficult to ascertain the structure of a protein in its active form. This is because proteins remain in their high energy, active conformations for too fleeting of time to obtain laboratory data through X-Ray crystallography or NMR.

- An alternative for obtaining the structures of high-energy protein conformations is through dynamics simulations
  - Unfortunately, many dynamics algorithms are comprised of too many degrees of freedom to run a simulation long enough to observe a change in protein conformation within a reasonable amount of computing time.

- The Weighted-Ensemble Brownian (WEB) Dynamics method proposed by Huber and Kim surpasses other algorithms in its ability to calculate the same length of simulation as other methods, but at only a fraction of the time.

- To fully understand the applicability of the WEB method, one must try it out for himself → My project.

versus
Current Research

- My research goal was to program a small-scale WEB Dynamics simulation in Perl and analyze the results
  - Quite the feat because I have never programmed before in my life
- The System
  - One-Dimensional
  - One Energy Barrier

\[ U(x) = E_b \left[ \left( \frac{x}{d} \right)^2 - 1 \right]^2 \]

where

\[ d = 1\text{Å} \quad E_b = 11k_B T \]
Methods

Standard Brownian Algorithm

\[ x_{i+1} = x_i + F_i \times (m \gamma)^{-1} \times \delta t + X_i(\delta t) \]

where:

\[ m = \frac{1k_BT S^2}{\AA^2} \]
\[ \delta t = 0.001\text{s} \]
\[ \gamma = 1\text{s}^{-1} \]

\[ F_i = -\frac{dU}{dx} \bigg|_{x_i} \quad X_i(\delta t) = \pm \sqrt{\frac{2k_BT \delta t}{m\gamma}} \]
Methods

- Run each particle through 100 iterations of a standard Brownian algorithm
  - Sort particles into proper bins
  - Modify (split/combine) particles so that each bin with any particles in it (active bin) will have 10 evenly-weighted particles in it after modification
  - Modify the particles’ weights in each active bin to ensure that the total weight of particles in the bin before modification is the same as the total weight in the bin after modification.
- Repeat this process 1000 times

\[
\frac{100 \text{ iterations}}{1 \text{ modification event}} \times 1,000 \text{ modification events} = 100,000 \text{ iterations}
\]

\[
100,000 \text{ iterations} \times \frac{.001 \text{ second}}{1 \text{ iterations}} = 100 \text{ second simulation}
\]
Methods

Initial System:
- 28 bins
- 10 evenly-weighted particles in bin 4
- 0 particles in other bins
- Bin division should follow particle movement

$$\sum_{i=1}^{N} w_i = 1$$

$N$ = total number of particles in the system

position = -1
weight = .1
Methods

First Modification Event (100 iterations {.1 seconds})

weight = .1
Methods

After Modification

\[ U(x) \]

weight = .03

weight = .04

weight = .01

weight = .02
Expected Results

- The Distribution of bin weight should follow the Boltzmann Distribution
  - Remember that the total weight of all the particles at all times is 1, which represents a probability distribution by bin
    \[
    P(x) = e^{\frac{-U(x)}{k_B T}}
    \]

To normalize:
\[
P(x) = \frac{e^{\frac{-U(x)}{k_B T}}}{Z}
\]

Partition Function:
\[
Z = \int_{-\infty}^{+\infty} e^{\frac{-U(x)}{k_B T}}
\]
Hurdles Along the Way

- I first tried a two-dimensional array, but had problem reassigning new values to the arrays representing the particles.
  - I then revised my approach to use two one dimensional arrays to define my particles.
- Sorting the particles was surprisingly easier than I expected; the only difficulty I had was indexing the particles so that the weight and position of each particle maintained correlation.
- I found splitting particle to ensure each bin that had any particles in it had 10 particles in much easier than combining particles while conserving probability.
- I learned that you cannot expect Perl to do anything, you have to tell it exactly what you want it and don’t want it to do.
Results

WEB

Standard Brownian

Bin Weights at $t = 0s$

Bin Weights at $t = 5s$
Results

WEB

Standard Brownian

Bin Weights at t = 20s

Bin Weights at t = 40s

Bin Weights at t = 20s

Bin Weights at t = 40s
Results

WEB

Standard Brownian

Bin Weights at t = 60s

Bin Weights at t = 80s

Bin Weights at t = 60s

Bin Weights at t = 80s
Results

WEB Standard Brownian

Bin Weights at $t = 100s$

As $t \to \infty$

9 Energy Barrier Crossing Events

$P(x)$

0.5

As $t \to \infty$

0 Energy Barrier Crossing Events

$-1 \times 10^{-10}$ $0$ $1 \times 10^{-10}$

$x$
Continuing Work

The WEB method can be expanded far beyond 1-D systems

2-D Twin Mountain Peak Energy Coordinate

Bottle Necking

Bins follow particle movement more directly
Conclusions

- The WEB method yields the same results as the standard Brownian method if permitted to run indefinitely → approaching Boltzmann Distribution
- The WEB method allows more particles to cross the energy barrier earlier on in the simulation → less computing time to determine a change in equilibrium
- Possible to use this method when we don’t know the energy coordinate
  - If we know the forces acting on a given protein, we can still run a WEB dynamics simulation
  - From the resulting probability distribution, we can determine the likelihood of a protein to remain in any given energy state
  - Much more cost effective and efficient than using NMR and X-Ray crystallography to isolate the numerous equilibrium structure of any given protein
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