# PHY407: Computational Physics Fall, 2015

Lecture 11: Random Processes, Part 2

### **Summary & Status**

- ☑ Weeks 1-3: Programming basics, numerical errors, numerical integration and differntiation.
- ☑ Weeks 4-5: Solving linear & nonlinear systems and Fourier transforms.
- ✓ Week 6: ODEs Part 1: RK4, Leapfrog, Verlet, adaptive time stepping; customizing python output
- ☑ Week 7: ODEs Part 2: Bulirsch-Stoer, Boundary Value Problems/shooting,
- ☑ Weeks 8-9: PDEs Part 1: Elliptic equation solvers, leapfrog time stepping, FTCS, Crank-Nicholson, Spectral Methods
- ☑ Week 10: Stochastic methods, Part 1: random numbers, monte carlo integration
- ☐ Week 11: Stochast methods, Part 2: recap, statistical mechanics ideas, simulated annealing approach to optimization.
- ☐ Week 12 (no lab): recap, discussion, extensions.

# PHY407: Computational Physics Fall, 2015

#### Lecture 11: Random methods/stochastics Part 2

- Recap from last week
- Statistical mechanics ideas
- The "Metropolis" algorithm
- Simulated Annealing

# Recap from last week

- Python's (pseudo) random number generator is the Mersenne Twister
- Transformation of distributions, e.g.

For uniformly distributed z, Poisson distribution  $p(x) = \mu e^{-\mu x}$  obtained from

$$x = -\frac{1}{\mu} \ln(1 - z)$$

- Using random number sequence to generate fractal structures.
- Monte Carlo integration.

# Monte Carlo: Importance Sampling

- Name two reasons we need Monte Carlo integration...
- Hit or Miss integration and mean value method have errors that vary as N<sup>-1/2</sup>
- Importance sampling chooses weights that favour largest integration values:

$$I = \int_{a}^{b} f(x) dx = \left\langle \frac{f(x)}{w(x)} \right\rangle_{w} \int_{a}^{b} w(x) dx$$

$$\left\langle \frac{f(x)}{w(x)} \right\rangle_{w} = \frac{\int_{a}^{b} \left( \frac{f(x)}{w(x)} \right) w(x) dx}{\int_{a}^{b} w(x) dx} \approx \frac{1}{N} \sum_{i=1}^{N} \frac{f(x_{i})}{w(x_{i})}, \quad x_{i} \text{ from } p(x) \propto w(x)$$

• For a system in equilibrium at temperature T, the probability of finding the system in any particular microstate 'i' is given by:

$$P(E_i) = \frac{\exp(-\beta E_i)}{Z}, \qquad Z = \sum_{i=1}^{ALL} \exp(-\beta E_i), \qquad \beta = \frac{1}{k_B T}$$

- where  $E_i$  is the energy of state i, and  $k_B$  is Boltzmann's constant.
- System undergoes transitions between microstates with probability of being in a particular microstate P(E<sub>i</sub>)
- To calculate a macroscopic property during a measurement, like the total energy, or magnetization, we need to average over the many microstates that the system visits during the measurement.

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Ε

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At given T, what determines the probability of a particular microstate?

E

What states are more likely? Those with high E or those with low E?

low E (e.g. E  $<< k_B T$ )  $\rightarrow$  exp(-E/ $k_B T$ ) bigger than state with higher E

# The Problem of Large Numbers

If we want to measure a quantity 'X' over the macrostate:

$$< X > = \sum_{i=1}^{ALL} X_i P(E_i)$$

where  $X_i$  is the value of the quantity in the  $i^{th}$  microstate and P is the probability of finding the system in that microstate.

- Typically, this sum has an enormous number of terms.
- Simple example: single mole of gas has  $10^{23}$  molecules. Assume each molecule had only 2 possible quantum states (gross underestimation), then the total number of microstates of the mole of gas is:  $3^{10^{23}}$  which is HUGE.

#### Monte Carlo Summation in Stat. Mech.

- Huge number of terms in sum 

  use Monte Carlo summation
- Randomly sample the terms in the sum and only use those as an estimate. Replace:

$$< X > = \sum_{i=1}^{ALL} X_i P(E_i)$$

with a sum over 'N' randomly sampled microstates:

$$\langle X \rangle = \frac{\sum_{k=1}^{N} X_k P(E_k)}{\sum_{k=1}^{N} P(E_k)}$$

 the denominator is needed to ensure the total probability over the sampled states is 1.

#### Monte Carlo Summation in Stat. Mech.

 It is only worth keeping the big terms in the sum if we want to compute this:

$$< X > = \sum_{i=1}^{ALL} X_i P(E_i)$$

• There are a lot of states with  $P(E_i)$  really small, with  $E_i >> k_B T$ , which is the case for most of the states:

$$P(E_i) = \frac{\exp(-E_i / k_B T)}{Z}$$

- To get a good estimate for the sum, need to preferentially choose terms where the integrand is non-negligible.
- So we should use importance sampling!

#### Importance Sampling

For an integral:

$$I = \int_a^b f(x) dx = \int_a^b w(x) \frac{f(x)}{w(x)} dx$$

$$\Rightarrow I = \left\langle \frac{f(y)}{w(y)} \right\rangle_{w} \int_{a}^{b} w(x) dx \approx \frac{1}{N} \sum_{k=1}^{N} \frac{f(y_{k})}{w(y_{k})} \int_{a}^{b} w(x) dx$$

• For a sum (what we have):

$$< X > = \sum_{i=1}^{ALL} X_i P(E_i) \approx \frac{1}{N} \sum_{k=1}^{N} \frac{X_k P(E_k)}{w_k} \sum_{i=1}^{ALL} w_i$$

What to choose for weight w to reduce the variance?

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- What to choose for weight w to reduce the variance?
- P(E<sub>i</sub>)! (Let's keep terms with with large P)

$$< X > \approx \frac{1}{N} \sum_{k=1}^{N} \frac{X_k P(E_k)}{P(E_k)} \sum_{i=1}^{ALL} P(E_i)$$

### Importance Sampling in Stat. Mech.

$$\langle X \rangle \approx \frac{1}{N} \sum_{k=1}^{N} \frac{X_k P(E_k)}{P(E_k)} \sum_{i=1}^{ALL} P(E_i)$$

- Two ways we can simplify this expression. What are they?
- (1)

• (2)

# Importance Sampling in Stat. Mech.

$$\langle X \rangle \approx \frac{1}{N} \sum_{k=1}^{N} \frac{X_k P(E_k)}{P(E_k)} \sum_{i=1}^{ALL} P(E_i)$$

- Two ways we can simplify this expression. What are they?
- (1) Cancel out the P's in the first sum:

$$\Rightarrow \langle X \rangle \approx \frac{1}{N} \sum_{k=1}^{N} X_k \sum_{i=1}^{ALL} P(E_i)$$

• (2) Sum of probabilities over all states is 1:

$$\Rightarrow < X > \approx \frac{1}{N} \sum_{k=1}^{N} X_k$$

## Importance Sampling in Stat. Mech.

$$\langle X \rangle = \frac{1}{N} \sum_{k=1}^{N} X_k$$

wow that's much simpler!

but remember: you are now choosing your k's from a non-uniform distribution p

our sampling points k come from:

$$p_{k} = \frac{w_{k}}{\sum_{i=1}^{ALL} w_{i}} = \frac{P(E_{k})}{\sum_{i=1}^{ALL} P(E_{i})} = P(E_{k})$$

• So essentially, to find the macroscopic <X> we randomly choose terms in the sum based on their Boltzmann probabilities.

#### Markov Chain Method

• One thing left to deal with: How do we pick states with probability  $P(E_k)$ ? Recall:

$$P(E_k) = \frac{\exp(-\beta E_k)}{Z}, \qquad Z = \sum_{i=1}^{ALL} \exp(-\beta E_i), \qquad \beta = \frac{1}{k_B T}$$

- To do it this way, we need Z, which is a sum over all states. But if we could do this, we wouldn't need Monte Carlo in the first place!
- Solution: Use the Markov Chain Method:
  - text goes into details on how to implement this method with a Metropolis algorithm
  - I will summarize it algorithmically...

# Markov Chain Method: The Algorithm

- 1. Choose a random starting state
- Calculate the energy of that state E<sub>i</sub>
- Choose a transition to a new state uniformly at random from allowed set
- 4. Calculate the energy of this new state
- 5. Calculate the acceptance probability for this transition:

$$P_a = \begin{cases} 1 & \text{if } E_j \leq E_i \\ \exp[-\beta(E_j - E_i)] & \text{if } E_j > E_i \end{cases}$$
 • Always accept a lower energy state. Sometimes accept a higher energy state. • Accept higher energy more often for higher temperature.

- higher temperature.
- Accept/reject the move according to the acceptance probability
- Measure the quantity you want 'X' in its current state & store it
- 8. Repeat from step 2.

Ok, lets try applying it...

# Ising Model

- Simple model of ferromagnetism, but demonstrates many of the physical characteristics of fancier models.
- Assume an object is made up of a collection of dipoles (e.g. electron spins) and the net magnetization is the sum of the magnetization of all the spins
- Ising model:
  - assume the spins can only point up or down.
  - the spins interact and favor parallel alignment of pairs of spins
  - the interactions are non-zero only between nearest neighbours (i.e. distance dependent).

# Ising Model

The macroscopic energy given by:

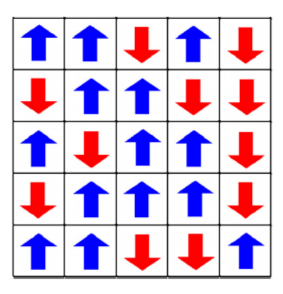
$$E = -J \sum_{\langle ij \rangle} s_i s_j$$

where s = 1 if spin is up & s = -1 if spin is down.

Notice that the lowest energy occurs if the spins all line up.

 Spins can randomly flip as the system visits a set of allowable states given its temperature. At any particular moment the system may

look like:



## Example: 1D Ising Model

- create array of dipoles, initial state: random spins
- Calculate energy & magnetization of state
- use Metropolis algorithm:
  - create new state: flip 1 spin randomly
  - calculate new total energy
  - calculate acceptance probability
  - decide whether to accept or reject new state
  - store 'new' energy & magnetization
  - repeat
- All right, lets do it...









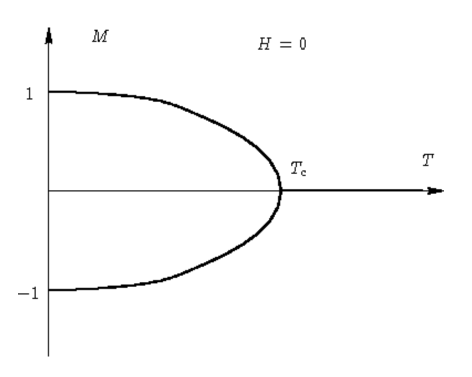


### 2-D Ising Model

- Goal: what happens at different T.
- What you should find:
- Low T: macroscopic magnetization occurs, but there are fluctuations in the microstates.
- Increasing T: lower macroscopic magnetization, more fluctuations
- Increase above a critical T (Curie temperature): no macroscopic magnetization, large fluctuations.

## Phase Transitions in the Ising Model

- Critical T: marks a phase transition.
- Above this value: no magnetization in system
- Below this value: magnetization.
- Note: there is no background magnetic field, so the system is symmetric about what sign of magnetization to acquire. This will depend strongly on the initial state of the system.
- Show simulations from python program...



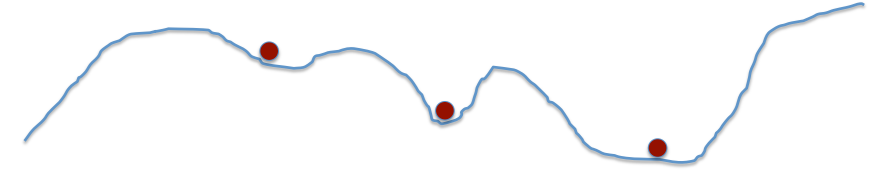
(for those who took nonlinear: physics: what kind of bifurcation is this?)

## Simulated Annealing

- Using Monte Carlo simulations to find GLOBAL minima/ maxima.
- In Ch. 6 we talked about ways of finding local mimima (e.g. golden ratio search).
- How it works: rewrite max/min problem as looking for a "ground state energy" of a system.
  - function f that you want the max/min: make this the energy function.
  - how could you find ground state: reduce temperature until you reach the ground state.

## Simulated Annealing

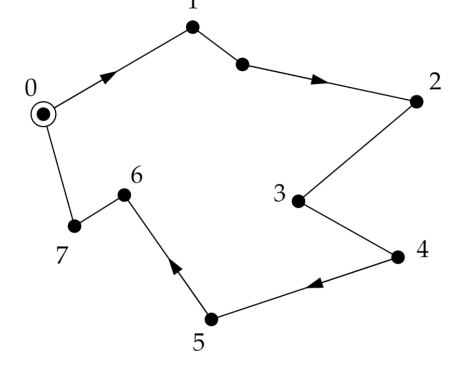
- Issue: if you reduce temperature too quickly: might get caught in a local min instead of the global min.
- Solution: reduce temperature slowly. This way system has time to explore many microstates and find a good approximation to the global minimum.
- Visual Analogy: particle in a bumpy potential. Too low energy: get stuck in nearest local minimum. Keep low energy but allow some random 'kicks' in energy: can kick out of local minimum and continue heading to global minimum.



# Example: Traveling Salesman

- Famous NP-hard problem: What is the shortest route to visit a given set of locations on a map?
- Want global minimum of distance
- start with random route, swap 2 cities, use Metropolis algorithm to determine whether to keep the swap
- 'energy' in this case is the total distance of the route

You will explore this problem using code from the book.



show salesman.py

#### Example: Minimization of a Function

- You will use simulated annealing to find the global minimum of a function f(x,y).
- Should be pretty straightforward to adapt from salesman.py.

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- The "Metropolis" algorithm
- Simulated Annealing