

# Monte Carlo of Molecular Systems

Chem 280

# Statistical Mechanics

the description of physical phenomena in terms of a statistical treatment of the behavior of large numbers of atoms or molecules, especially with regard to the distribution of energy among them.

--Oxford Languages

# Monte Carlo of Molecular Systems

According to statistical mechanics

We can use MC to evaluate this integral!

$$\langle Q \rangle = \int_V Q(r^N) \rho(r^N) dr^N$$

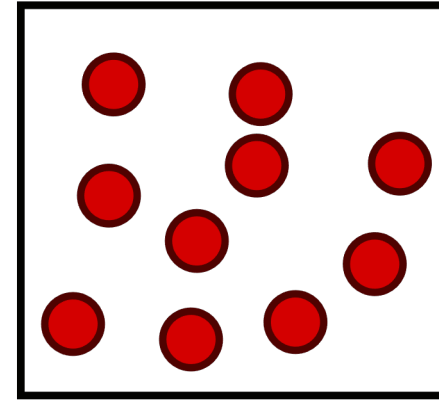
- $Q$  quantity which depends on atomic coordinates ( $r^N$ )
- $\langle Q \rangle$  average value of quantity  $Q$  (square brackets denote average)
- $r^N$  atomic coordinates of  $N$  atoms.
- $\rho(r^N)$  probability density based on thermodynamic properties (beyond scope of this course)

# Monte Carlo of Molecular Systems

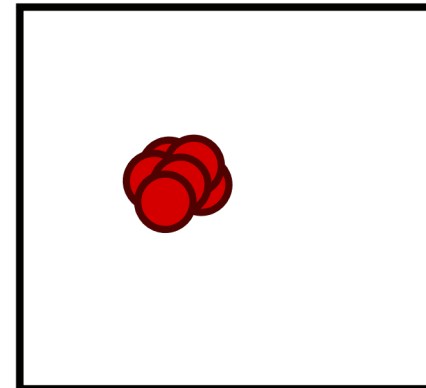
In order to evaluate this integral we have some special considerations

$$\langle Q \rangle = \int_V Q(r^N) \rho(r^N) dr^N$$

Because we have so many possible states, it is not effective to sample points with a uniform distribution. We want to sample configurations which are likely to occur.



Consider our 10 particles in a box.



This configuration (particles stacked) is high energy and not likely to occur

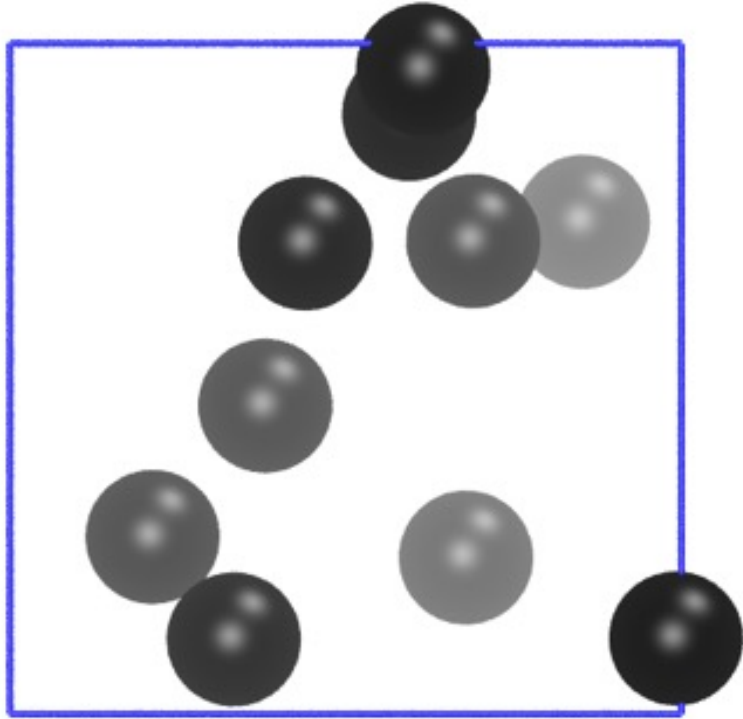
# Importance Sampling

- No longer using a uniform distribution for coordinate generation.
- Instead, generate configurations with distribution -  $\rho(\mathbf{r}^N)$  - the probability density based on thermodynamic properties.

Then, we can evaluate the integral as the average of the generated configurations:

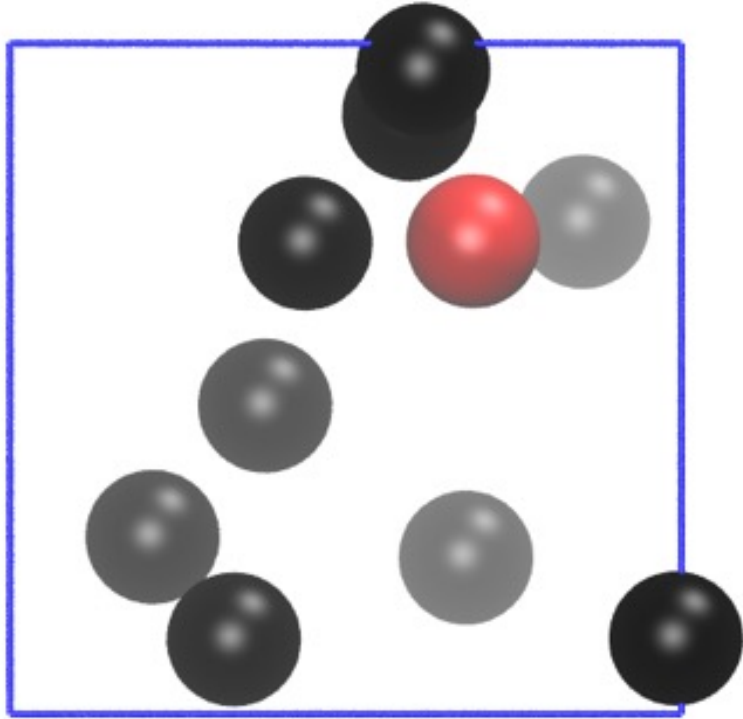
$$\langle Q \rangle = \frac{1}{N} \sum_{i=1}^N Q(\mathbf{r}_i^N)$$

# The Metropolis Monte Carlo Recipe



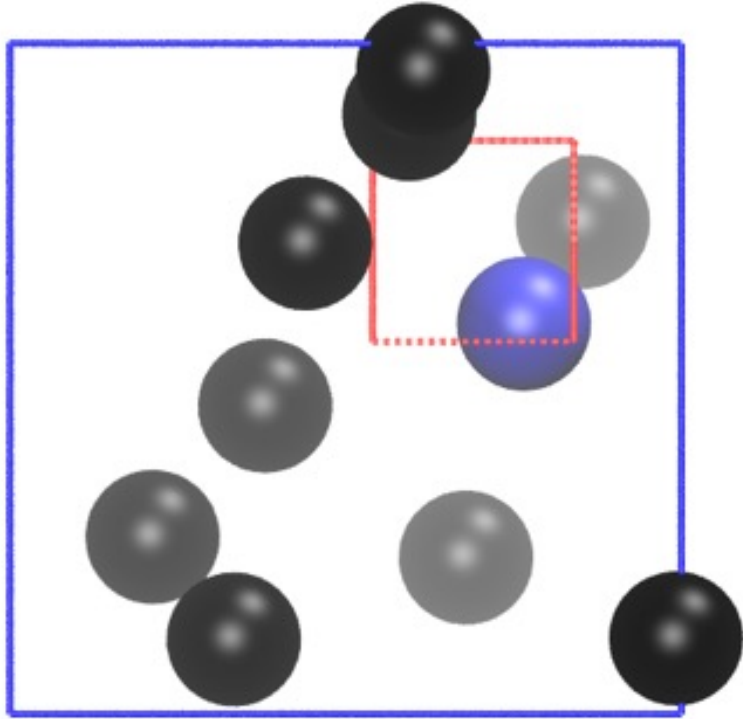
Generate an initial state  $m$   
and calculate its energy.

# The Metropolis Monte Carlo Recipe



Choose an atom with  
uniform probability

# The Metropolis Monte Carlo Recipe

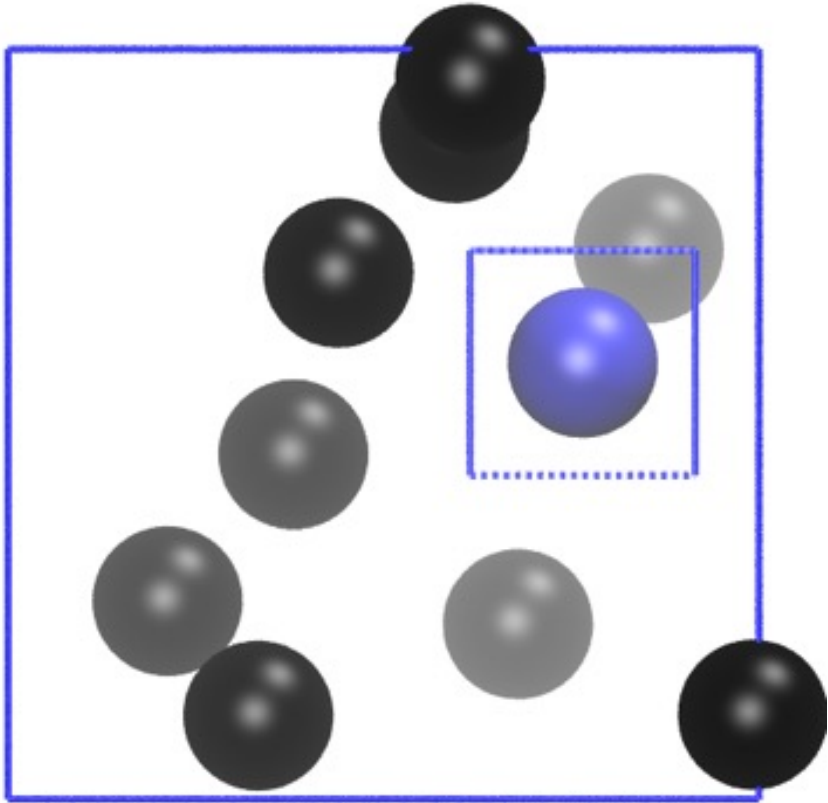


Attempt a random translation within a maximum distance.

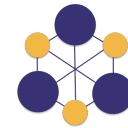
Calculate the energy of the new state,  $n$ .



# The Metropolis Monte Carlo Recipe



Accept or reject new  
state according to  
the Metropolis  
criterion



# The Metropolis Criterion

Accept move based on the energy change resulting from moving the particle and system temperature.

$$P_{acc}(m \rightarrow n) = \min[1, e^{-\Delta U/T}]$$

This means we will always accept moves which result in a decrease in energy ( $-\Delta U$ ), and sometimes accept moves which are zero or positive.

In practice, we will generate a random number on the range zero to 1. If our calculated  $P_{acc}$  is greater than our generated number, we accept the configuration.

# Reduced Units

For Argon,

$$\varepsilon = 120 K (k_B) = 1.68 \times 10^{-21} J \text{ and } \sigma = 3.4 \times 10^{-10} \text{ meters}$$

These are really inconvenient numbers!

We will normalize our energy by  $\varepsilon$  and our distances by  $\sigma$ .

$$U^*(r) = \frac{U(r)}{\varepsilon}$$

$$r^* = \frac{r}{\sigma}$$

$$U^*(r^*) = 4 \left[ \left( \frac{1}{r^*} \right)^{12} - \left( \frac{1}{r^*} \right)^6 \right]$$

This will make  
 $U^*(r^*)$  be on  
the order of 1.