

Simulation of Polymers

- Physics of polymers
- MD of polymers?
- MC methods
- Lattice models
- Reptation Monte Carlo
- Rosenbluth growth methods
- Pivot method

"Computer simulation methods for polymer physics"

Kurt Kramer in *MC and MD in Condensed Matter Systems*

"MC and MD simulations in polymer science"

K. Binder editor, Oxford, 1995.

4/8/2013

1

Time estimate for MD

- Time scales
 - Local oscillations are 10^{-13} s so time step is 10^{-14} s
 - Important motions in polymers take seconds or hours (real time) requiring 10^{14} to 10^{18} steps!
 - A system of 100 chains of 50 monomers (20,000 particles) takes about 1step/sec for 10^{-4} s (real time) would take about 10^{10} secs or 300 years!
- Distance scales
 - Local effects are order 1A.
 - Volume of cell is $(100\text{A})^3$.
- Solvent is important. Hydrodynamic effects dominate.
- Conclusion: You need to make a simplified model of the polymer to do research in this area unless you have a much faster computer.

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2

Polymer Hamiltonian

- **Self-avoiding random walk. (SAW)**
Consider a simple lattice and take a random walk on the lattice--one which only visit each site once.
- **Bead spring model**
 - Bonding interaction holds the chain together. Key feature of polymer. *A bead does not represent an atom, but a blob--a section of the chain.*
 - Non-bonded excluded volume interaction (LJ)

$$V = \sum_{i < j} \phi(r_{ij}) + \sum_i \gamma(|r_i - r_{i+1}|)$$

Modified harmonic potential (with $k = 20, \sigma = 1.95\sigma_{HS}$)

$$\gamma_h = \frac{1}{2}k(r - r_0)^2 \quad \gamma_{hl} = -\frac{1}{2}k\sigma \ln[1 - (r / \sigma)^2]$$

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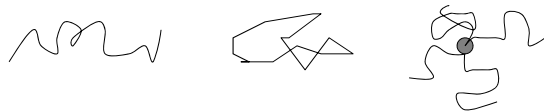
3

Polymer Phases

- For a repulsive interaction--the chains stretch out, swell.
- Characterize size by mean square end-end distance.

$$\langle (r_n - r_0)^2 \rangle \propto N^{2\nu}$$

$$\nu \sim 0.588 \text{ SAW} \quad \text{or } 0.5 \text{ RW.}$$
- **This means MD will be very slow. Relaxation time = $N^{2.2}$.**
- As attractive interaction are added in
 - at some point the polymers collapse. (*Theta point* collapse.)
 - Right at collapse point--walks are *uncorrelated* random walks.
 - This is a type of phase transition.
- Big question: **how does the dynamics scale with the length of the chain-entanglement?**
- Other topologies for polymers: linear, rings, stars,

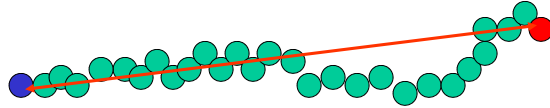


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4

Polymer Reptation (slithering snake)

- Polymers move very slowly because of entanglement.
- Local MC just as slow as MD.
- A good algorithm is "reptation."
 - *Cut off one end and stick onto the other end.*
 - *Choose end at random or "bounce" with rejection.*



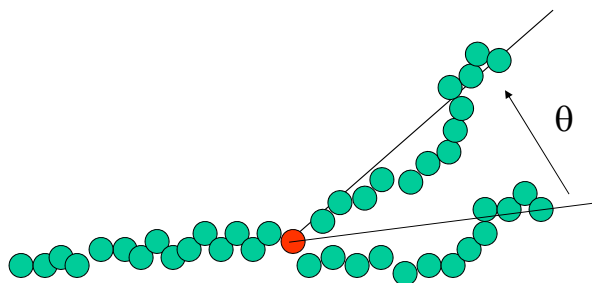
- Sample directly the bonding interaction
- Acceptance probability is change in non-bonding potential.
- Simple moves go quickly through polymer space.
 - But Ergodic? Not always (what if both ends get trapped?)
- Decorrelation time is $O(N^2)$. Works for many chains.
- Completely unphysical dynamics or is it?
 - This may be how entangled polymers actually move. (theory of Degennes)

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7

Pivot algorithm

- Take a polymer. Pick an atom at random.
- Rotate one segment with respect to the pivot point a random angle θ .
- Accept or reject.
- Most efficient method for a single chain. Exponent of relaxation of end-end distance is $N^{0.2}$

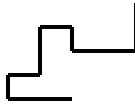


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8

Lattice model for polymers

- Maybe we can speed up the algorithm by forcing the polymer to lie on a lattice.
- SAW = "self-avoiding random walk": a walk on a lattice with N steps which cannot visit a site more than once.



- Partition function=sum over all such possible walks.
- Monte Carlo=sample the distribution of the walks and take averages such as end-end distribution.
- You can also put a "non-bonded" interaction to make polymer collapse.

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9

How to move polymers

- Growth
- Reptation
- Crankshaft moves



- If move is allowed, accept it.
- Pivot moves
-

Ergodic questions arise:

Can you go everywhere in chain space?

Make a mixture of moves.

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10

Growth algorithms

CBMC = Configurational Bias MC/Rosenbluth
Chapters 11,13 FS

- Simply grow polymer, stopping when you get any overlap.
- Use importance sampling to direct the walk in favorable directions.

$$W_n = W_{n-1} \left[\frac{q_i}{q} \right] \quad \langle R^2 \rangle = \frac{\sum W_j R_j^2}{W_i}$$

q_i = # of open moves

- Problem: **can you get really long polymers?**
- Use branching when weights fluctuate too much.
- Easily generalized to continuum models.
- In CBMC we grow a new section and accept or reject it

$$acc. prob. = \min \left[1, \frac{W_{new}}{W_{old}} \right]$$