Polymer chains

Ludwig Rasmijn 4106644

Sebastiaan Lokhorst 4005058 Shang-Jen Wang 4215974

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Abstract

The behavior of polymer chains in 2D is investigated by simulating a random walk using two different methods. Both the Rosenbluth and the Pruning-Enriched Rosenbluth Method (PERM) are used, with six choices for successive beads, which have a probability based on the energy of the new bead, which is modeled with the Lennard-Jones potential. The end-to-end distance R as a function of the length of the polymer, expressed in the number of beads N, is studied for both methods. It is concluded that they are related by $R \propto (N-1)^{0.75}$, which is in accordance with literature such as Thijssen[1].

1 Introduction

Rosenbluth algorithm and optimizations are used to simulate polymer chains.

2 Methods

The polymer model is a self-avoiding walk in 2 dimensions, where each effective monomer is a fixed distance from the nearest neighbour. The interaction between the beads is modeled by the Lennard-Jones potential:

$$V_{LJ} = 4\epsilon \left[\left(\frac{\sigma}{r} \right)^{12} - \left(\frac{\sigma}{r} \right)^{6} \right]. \tag{1}$$

Where $\epsilon=0.25$ (natural units), and $\sigma=0.8$, given that the distance between monomers is equal to 1.

2.1 Rosenbluth algorithm

The Rosenbluth algorithm is a Monte-Carlo method to generate polymers, which will avoid high-energy conformations when adding new beads to the polymer.[2] This method uses six evenly spaced $(2\pi/6)$ discrete angles with a random offset for which the new bead can be added to the polymer, as shown in fig. 1. It will avoid any improbable conformations by adding the next bead with a probability proportional to $\exp(-E(\theta)/(k_BT))$, where $E(\theta)$ is the interaction energy of the new bead with the rest of the polymer, and θ is the angle between the new bead and the previous two beads. The weight for each angle θ_i is calculated with:

$$w_j^{(l)} = \exp(-E(\theta_j)/(k_B T)),$$
 (2)

where j is the number of the angle and l is the bead that is being added. The sum of all the weights can be expressed as:

$$W^{(l)} = \sum_{j} \exp(-E(\theta_j)/(k_B T)).$$
 (3)

So angle j is accepted with a probability $w_i^{(l)}/W^{(l)}$.

The weight of the total polymer is given by:

$$PolWeight = \prod_{l} W^{l}.$$
 (4)

2.2 Pruned-enriched Rosenbluth method

A method which suppresses the high energy configuration better than the Rosenbluth method is the pruned-enriched Rosenbluth method (PERM)

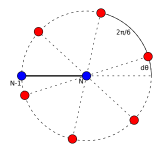


Figure 1: Choosing bead N+1 out of one of the six possible angles θ_i

by Grassberger [3]. This method removes configurations with a high energy and low weight and multiplies configurations with a low energy and high weight. This way, the high-weight and thus more important configurations, are studied more closely.

Two thresholds UpLim and LowLim are used to define 'low' and 'high' weights. If at any polymer length the PolWeight > UpLim, then two members of this polymer are created when adding the next bead and each are given a PolWeight which half the PolWeight of the original polymer. This is called 'enriching'. If at any polymer length the PolWeight < LowLim, then the polymer will be removed with a probability of 1/2. If the polymer is not removed, then the PolWeight is multiplied by 2. This will make sure that the distribution does not change. This is called 'pruning'.

The choice of UpLim and LowLim depends on the average weight AvWeight at polymer length L. The average weight is updated for every polymer that reaches that length. UpLim and LowLim are expressed as a ratio of the average weight and the weight Weight3 corresponding to the shortest length (3 beads):

$$UpLim = \alpha \cdot AvWeight/Weight3, \qquad (5)$$

$$\texttt{LowLim} = \alpha \cdot \texttt{AvWeight/Weight3}. \tag{6}$$

A good value for α for UpLim is 2 and for LowLim 1.2. It is possible that α is dependent on L, but this dependency can be removed by multiplying the weight of the polymer by a constant at each step. This constant should be near $1/(0.75N_{\theta})$, where N_{θ} is the number of possible angles.

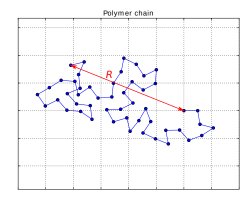


Figure 2: End-to-end distance R for a polymer of length N=50

3 Results and Discussion

The polymer simulation was used to study the end-to-end distance of a polymer R, illustrated in fig. 2.

An average value for R^2 can be obtained by simulating multiple polymers and then averaging them with the weight of the polymers.

The end-to-end distance R of the polymer should scale with:

$$R \propto N_s^{\ \nu},$$
 (7)

where N_s is the number of segments, and $\nu = 0.75$ in two dimensions.[1]

This means that the results should fit:

$$R^2 = a(N-1)^{1.5}, (8)$$

Where a is a scaling constant.

The square end-to-end distance R^2 , determined with the regular Rosenbluth algorithm is shown in fig. 3. Note that the population stays constant but the error for large N is quite large, even for such an high population.

The same \mathbb{R}^2 , but now determined with the PERM algorithm, is shown in fig. 4. Note that while the population is *smaller* than for the regular Rosenbluth method, the error for large N is much smaller. This is because low-weight (and thus irrelevant) configuration are suppressed, and the computation focuses on the high-weight (relevant) population.

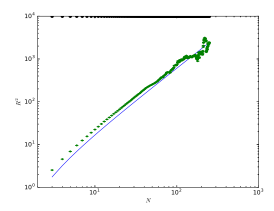


Figure 3: The average R^2 as a function of the number of beads N. Fitted to the function $R^2 = 0.62(N-1)^{1.5}$. The black dots represent the population size, which is constant at 10,000

Because in the PERM case the computation is focused on the relevant polymers, it is much more efficient. The computation time scales with $\sum_{N} \text{population}(N)$. When comparing the population of the regular Rosenbluth and the PERM algorithm, they both are relatively constant. However the error in the case of the PERM algorithm is much better, even with a population which is ten times as small.

4 Conclusion

In this study, polymer chains were simulated using the Rosenbluth and Pruning-Enriched Rosenbluth Method (PERM). Polymers were grown to a length of up to 250 beads. The end-to-end distance was investigated and found to be in accordance with literature, namely obeying the relationship $R^2 = a(N-1)^{1.5}$ with $a \approx 0.6$. It was found that the PERM method is superior in finding high-probability configurations and thus requiring a smaller population to achieve a more precise result.

References

[1] J.M. Thijssen (2007) Computational Physics.

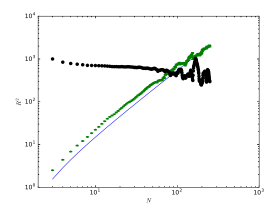


Figure 4:

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