

Polymer Chain Dynamics Simulation Report

ZHA Chenyu

May 12, 2015

1 The random walk model:the freely jointed chain

Consider a linear polymer to be a freely-jointed chain with N beads, length of each bond is b , that occupy zero volume. The path of the chains is like a 'random walk' in three dimensions, limited only by the constraint that each segment must be joined to its neighbors.

Consider the 'end to end' vector \mathbf{R} joining one end of the polymer to the other, the average value $\langle \mathbf{R} \rangle$ is zero, since the probability of \mathbf{R} equals $-\mathbf{R}$. Therefore we will calculate $\langle \mathbf{R}^2 \rangle$

$$\langle \mathbf{R}^2 \rangle = \sum_{n=1}^N \sum_{m=1}^N \langle \mathbf{r}_n \cdot \mathbf{r}_m \rangle \quad (1)$$

We consider that there is no correlation between bead n and m , therefore we find :

$$\langle \mathbf{R}^2 \rangle = \sum_{n=1}^N \langle \mathbf{r}_n^2 \rangle = Nb^2 \quad (2)$$

The probability distribution of \mathbf{R} is:

$$P(\mathbf{R}, N) = \left(\frac{3}{2\pi Nb^2} \right)^{3/2} \exp\left(-\frac{3\mathbf{R}^2}{2Nb^2}\right) \quad (3)$$

The probability distribution function of \mathbf{R} obeys the Gaussian distribution. The position of the beads after each step will satisfy the diffusion equation :

$$\mathbf{R}(t + \Delta t) = \mathbf{R}(t) + \sqrt{2D\Delta t} \mathbf{g}(t) \quad (4)$$

$\mathbf{g}(t)$ is normally distributed random noise.

1.1 Random walk Simulation

```
%This program is used to simulate the brownian motion
classdef FirstTry<handle

properties
    dimension%dimension=1,2 or 3
    numParticles%number of particles in polymers;
    dt%pas de temps
    numSteps% number of motion
    diffusionConst %constante diffusion
    paths %the paths of polymer;
end

methods

%class constructor
function obj=FirstTry(dimension,numParticles,dt,diffusionConst,numSteps)
```

```

obj.dimension=dimension;
obj.numParticles=numParticles;
obj.dt=dt;
obj.numSteps=numSteps;
obj.diffusionConst=diffusionConst;
obj.paths = zeros(obj.numParticles,3,obj.numSteps);
end

function Calculate(obj)

for j=1
noise = [zeros(1,obj.dimension);...
sqrt(2*obj.diffusionConst*obj.dt)*randn(obj.numParticles-1,obj.dimension)];
obj.paths(:,1:obj.dimension,j)=cumsum(noise);

end

for j=2:obj.numSteps
noise = [sqrt(2*obj.diffusionConst*obj.dt)*randn(obj.numParticles,obj.dimension)];
obj.paths(:,1:obj.dimension,j)=obj.paths(:,1:obj.dimension,j-1)+noise;

end

end %'random walk 'simulation

function Plot(obj)
f=figure;
b =[-20 20];
a= axes('Parent',f,'NextPlot','replaceChildren','XLim',b,'YLim',b,'ZLim',b);
c=rand(1,3);

x = obj.paths(:,1,1);
y = obj.paths(:,2,1);
z = obj.paths(:,3,1);
% end
l=line('XData',x,'YData',y,'ZData',z,'Color',c,'linestyle','-','Marker','o','markersize',10,'Parent',f);
for i=2:obj.numSteps

set(l,'XData',obj.paths(:,1,i),'YData',obj.paths(:,2,i),'ZData',obj.paths(:,3,i));

pause(1)

drawnow

end
end

end
end

```

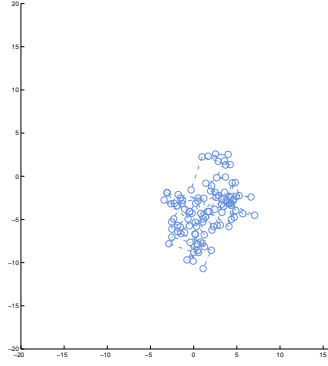


Figure 1: initial position

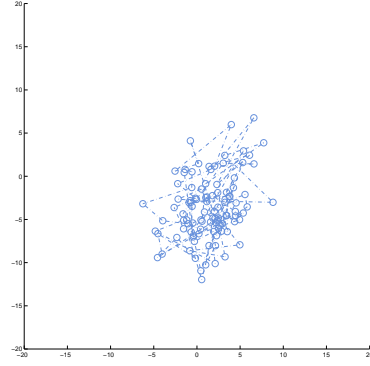


Figure 2: final position

1.2 Probability distribution function of R

In order to verify that the PDF of R is Gaussian, we calculate the 'end to end distance' R for each simulation, then we plot it with histogram by coordinate (x, y, z) respectively and compare with the probability distribution function of R in theory.

```
%this program is used to simulate the probability distribution of Gaussian
classdef Idealchain<handle

properties
    dimension%dimension=3
    numParticles%number of particles in polymers;
    dt%pas de temps
    numSteps% number of motion
    diffusionConst %constante diffusion
    paths %the paths of polymer;
    endToEndDist %end to end distance
    simulation %number of simulations
end

methods

%class constructor
function obj=Idealchain(dimension,numParticles,dt,diffusionConst,numSteps,simulation)
    obj.dimension=dimension;
    obj.numParticles=numParticles;
    obj.dt=dt;
    obj.numSteps=numSteps;
    obj.diffusionConst=diffusionConst;
    obj.simulation=simulation;
    obj.paths = zeros(2,3,2);
    obj.endToEndDist=zeros(obj.simulation,3);
end
```

```

function Calculate(obj)
for s=1:obj.simulation
%step 1:connect the chain;
obj.paths(1,1:obj.dimension,1)=[0 0 0];%position of first beed;
obj.paths(2,1:obj.dimension,1)=obj.paths(1,1:obj.dimension,1);
noise=sqrt(2*obj.diffusionConst*obj.dt)*randn(1,obj.dimension);
for i=1:obj.numParticles
obj.paths(2,1:obj.dimension,1)=obj.paths(2,1:obj.dimension,1)+noise;%position of last beed;
end

%step 2:end :the position of beed 1 and beed end varity by time;
for j=2:obj.numSteps
obj.paths(1,1:obj.dimension,2)=obj.paths(1,1:obj.dimension,1)+noise;
obj.paths(2,1:obj.dimension,2)=obj.paths(2,1:obj.dimension,1)+noise;
obj.paths(1,1:obj.dimension,1)=obj.paths(1,1:obj.dimension,2);
obj.paths(2,1:obj.dimension,1)=obj.paths(2,1:obj.dimension,2);
end
obj.endToEndDist(s,:)=obj.paths(2,:,2)-obj.paths(1,:,2);

end

end

function Plot(obj)
%
%         figure(2);
%         plot(obj.endToEndDist);

%
%         figure(3)
%         [h,bins]= hist(obj.endToEndDist(:,1),50);
%         bar(bins,h);
%

f=@(N,R,b)(sqrt(1/(2*pi*N*b^2))*exp(-(sum(R.^2,2))./(2*N*b^2)));%PDF
subplot(2,2,1)
[h, bins]=hist(obj.endToEndDist(:,1),50); h= h./sum(h); bar(bins,h),...
hold on,
plot(bins,f(obj.numParticles,[bins'],1),'r')
title('PDF on x')

subplot(2,2,2)
[h, bins]=hist(obj.endToEndDist(:,2),50); h= h./sum(h);bar(bins,h),...
hold on, plot(bins,f(obj.numParticles,[bins'],1),'r')
title('PDF on y')

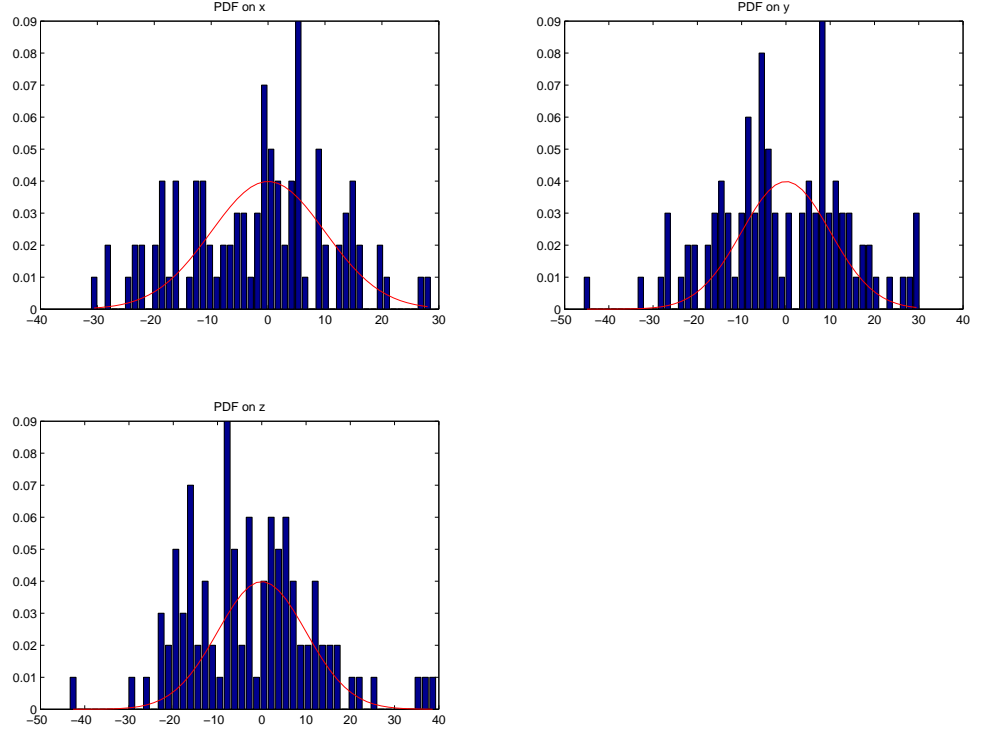
subplot(2,2,3)
[h, bins]=hist(obj.endToEndDist(:,3),50); h= h./sum(h); bar(bins,h),...
hold on, plot(bins,f(obj.numParticles,[bins'],1),'r')
title('PDF on z')
%plot(3/(2*pi*obj.numParticles*1)^1.5*exp(3*obj.R.^2/(2*obj.numParticles*1)))

end

end

end

```



2 The bead-spring model

The bead-spring model is also called Rouse Model .In this model,the single chain diffusion is represented by Brownian motion ,there is no exclude volume interactions between the beads and each bead experience a drag force proportional to their velocity ,then the position of the beads will satisfy the Langevin equation :

$$\frac{d\mathbf{R}_n}{dt} = \frac{k}{\xi}(\mathbf{R}_{n+1} + \mathbf{R}_{n-1} - 2\mathbf{R}_n) + \mathbf{g}_n, \forall n \in [1, 2 \dots N - 1] \quad (5)$$

For the bead 0 and N ,we have :

$$\begin{cases} \frac{d\mathbf{R}_0}{dt} = \frac{k}{\xi}(\mathbf{R}_1 - \mathbf{R}_0) + \mathbf{g}_n \\ \frac{d\mathbf{R}_n}{dt} = \frac{k}{\xi}(\mathbf{R}_{n-1} - \mathbf{R}_n) + \mathbf{g}_n \end{cases}$$

where ξ is friction coefficient, k is spring constant.

2.1 Rouse Model Simulation

```
%This program is used to simulate the spring-bead model
classdef RouseModel<handle
properties
    dimension%dimension=1,2 or 3
    numParticles%number of particles in polymers;
    dt%pas de temps
    numSteps% number of motion
    numSimulations%number of simulations;
    diffusionConst %constante diffusion
    paths %the paths of polymer;
    frictionCoefficient;%the frictionCoefficient of a bead;
    MetBeedNum; %the number of the beed which have met with the first and last beed;
    connectedBeads %an n by two array with pair wise bead numbers to connect
    fixedBeads %numBeads of beads that do not move
    b %length between 2 beads
end

methods

function obj=RouseModel(dimension,numParticles,dt,diffusionConst,...
numSteps,frictionCoefficient,connectedBeads, fixedBeads,b)

    obj.dimension = dimension;
    obj.numParticles = numParticles;
    obj.dt = dt;
    obj.numSteps = numSteps;
    obj.diffusionConst = diffusionConst;
    obj.paths = zeros(obj.numParticles,3,obj.numSteps);
    obj.frictionCoefficient = frictionCoefficient;
    obj.fixedBeads= fixedBeads;
    obj.connectedBeads = connectedBeads ;
    obj.b=b;
end

function Simulation(obj)

% set initial position
for j=1
    noise = [zeros(1,obj.dimension);...
    sqrt(2*obj.diffusionConst*obj.dt)*randn(obj.numParticles-1,obj.dimension)];
    obj.paths(:,1:obj.dimension,j)=cumsum(noise);

end

a = obj.dimension*obj.diffusionConst/obj.b^2;
R = RouseMatrix(obj.numParticles, obj.connectedBeads, obj.fixedBeads);

for j=2:obj.numSteps

    noiseSingle = sqrt(2*obj.diffusionConst*obj.dt)*randn(obj.numParticles,obj.dimension);
    %
    obj.paths(:,j)=obj.paths(:,j-1)+obj.dt*(a*(obj.paths(2,:,j-1)-obj.paths(1
```

```

% zero out noise for fixed particles
noiseSingle(obj.fixedBeads,:) = 0;

obj.paths(:,1:3,j) = -a*R*obj.paths(:,1:3,j-1)*obj.dt+noiseSingle +obj.paths(:,1:3,j-1);

end

end

function Plot(obj)
f=figure;
b=[-10 10];
d= axes('Parent',f,'NextPlot','replaceChildren','XLim',b,'YLim',b,'ZLim',b);
daspect([1 1 1]);
cameratoolbar
c=rand(1,3);

x = obj.paths(:,1,1);
y = obj.paths(:,2,1);
z = obj.paths(:,3,1);

l=line('XData',x,'YData',y,'ZData',z,'Color',c,'linestyle','-','Marker','o','markersize',10,'P
for i=2:obj.numSteps

set(l,'XData',obj.paths(:,1,i),'YData',obj.paths(:,2,i),'ZData',obj.paths(:,3,i));

pause(0.2)

drawnow
end

end

end
end
end

```