# Ch/ChE 164 Winter 2024

Midterm

Due Date: Wednesday, February 7, 2024

Midterm Policy

- This is an open-book exam. You may look at your notes, your homework, the course notes, and the books listed on the course syllabus.
- You may not refer to homework or exams from previous years. You may not consult the internet outside of the course canvas page.
- You may take as long as you need before the due date to work on the exam.
- You may not consult with one another.
- Please write clearly and show all steps; points will be deducted if the midterm is unclear or illegible.
- Midterm must be turned in by the end of day on the due date. Late midterm will be penalized 10% per day unless prior arrangements have been made with the instructor.

Some identities that may or may not be useful (not necessarily exhaustive)...

$$(x+y)^N = \sum_{n=0}^N \binom{N}{n} x^n y^{N-n}$$

$$(1+x)^N = \sum_{n=0}^N \binom{N}{n} x^n = \sum_{n=0}^N \binom{N}{n} x^n (1)^{N-n}$$

$$\frac{1}{1-x} = \sum_{n=0}^\infty x^n \text{ for } |x| < 1$$

$$\cosh(x) = \frac{e^x + e^{-x}}{2} = \frac{d}{dx} \sinh(x)$$

$$\sinh(x) = \frac{e^x - e^{-x}}{2} = \frac{d}{dx} \cosh(x)$$

$$\tanh(x) = \frac{\sinh(x)}{\cosh(x)}$$

## 1

1. Cross-linking polymer chain (30 pts) Consider polymer chains, each with N monomers, all of which can reversibly associate with any other monomer. For simplicity, let us consider a single polymer chain at temperature T.

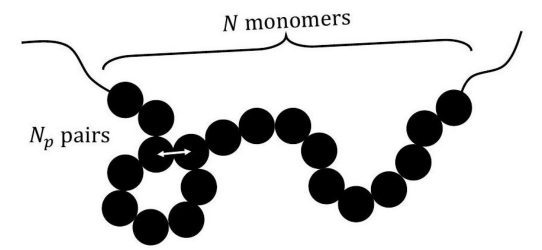


Figure 1: Visualisation of a polymer chain with N monomers forming a single, reversible pair.

(a) (3 pts) How many ways can you select monomers from the chain to form  $N_p$  pairs?

### 1.1

For this part, there will be two variants: one where N is even and one where N is odd. When even, this is:

$$\binom{N}{2N_p} \tag{1}$$

When odd, this is:

$$\binom{N-1}{2N_p} \tag{2}$$

but in the thermodynamic limit, this can well be approximated by the even case.

(b) (5 pts) From these monomers you've selected in 1a, how many ways can you form  $N_p$  pairs (it may be easier to first consider the case where you can form 2 pairs from 4 monomers)?

### 1.2

We will start by considering the simple case of forming 2 pairs from 4 monomers: We can pair the first one with the second one, third one, or fourth one. Because we only have two monomers left unpaired, there are no extra combinatorics associated with this. Now let us consider the case of 6 monomers: Once we establish one pair, the problem reduces to our initial one of forming 2 pairs from 4 monomers. This is a recursive problem. So, for the first pair there are 5 options, then there are only 3 options for the 4 remaining monomers, as discussed previously, and only 1 option for the last pair. So there are  $(2N_p - 1)!!$  ways to form  $N_p$  pairs from  $N_p$  monomers.

Note: You may need to use  $(N-1)!! = (N-1) \times (N-3) \times \dots$  This can be simplified using:  $(N-1)!! = N! / \lceil (N/2)! 2^{N/2} \rceil$ .

(c) (6 pts) The product from  $\mathbf{1a}$  and  $\mathbf{1b}$  will give you the microcanonical partition function for a system with  $N_p$  pairs as:

$$\Omega\left(N,N_{p}\right) = \frac{N!}{(N-2N_{p})!N_{p}!2^{N_{p}}}$$

If the energetic gain from forming a single pair is  $-\epsilon$ , write out the canonical partition function, Q(N,T), for this system. You should end up with a sum over all possible values of  $N_p$ .

### 1.3

The canonical partition function would be given by:

$$Q(N,T) = \sum_{N_p=0}^{N/2} \Omega(N, N_p) e^{-\beta \epsilon N_p}$$
(3)

Inserting the expression for  $\Omega(N, N_p)$ , we get:

$$Q(N,T) = \sum_{N_p=0}^{N/2} \frac{N!}{(N-2N_p)! N_p! 2^{N_p}} e^{-\beta \epsilon N_p}$$
 (4)

(d) (10 pts) Find the most-probable term in this sum, and show that the corresponding value for the fraction of monomers associated  $(p = 2N_p/N)$  is given by:

$$p = 1 + \frac{\frac{\exp(-\beta\epsilon)}{N} - \sqrt{\left(2 + \frac{\exp(-\beta\epsilon)}{N}\right)^2 - 4}}{2}$$

### 1.4

An arbitrary term in the sum is given by:

$$t_{N_p} = \frac{N!}{(N - 2N_p)! N_p! 2^{N_p}} e^{-\beta \epsilon N_p}$$
 (5)

We can find the most-probable term by taking the derivative of  $t_{N_p}$  with respect to  $N_p$  and setting it equal to zero. Or alternatively, we can choose to maximize the logarithm of  $t_{N_p}$ , which is equivalent, so the first derivative of the logarithm of  $t_{N_p}$  with respect to  $N_p$  is zero:

$$\frac{d}{dN_p}\ln(t_{N_p}) = 0 (6)$$

The logarithm of  $t_{N_n}$  is given by:

$$\ln(t_{N_p}) = \ln(N!) - \ln((N - 2N_p)!) - \ln(N_p!) - N_p \ln(2) - \beta \epsilon N_p$$
 (7)

Using Stirling's approximation, we can simplify the logarithm of  $t_{N_n}$  to:

$$\ln(t_{N_p}) \approx N \ln(N) - (N - 2N_p) \ln(N - 2N_p) - N_p \ln(N_p) - N_p \ln(2) - \beta \epsilon N_p$$
(8)

We can now take the first derivative of this with respect to  $N_p$  and set it equal to zero:

$$0 = -\beta \epsilon - \log(N_p) + 2\log(N - 2N_p) - \log(2) + 1 \tag{9}$$

Bring the non-logarithmic terms to the left-hand side:

$$\beta \epsilon - 1 = -\log(N_p) + 2\log(N - 2N_p) - \log(2) \tag{10}$$

Exponentiate both sides:

$$\exp\{(\beta \epsilon - 1)\} = \frac{(N - 2N_p)^2}{2N_p}$$
 (11)

Solving for  $N_p$ :

$$N_p = \frac{2eN - \sqrt{(4eN + e^{\beta\epsilon})e^{\beta\epsilon} + e^{\beta\epsilon}}}{4e}$$
 (12)

```
1 from sympy import symbols, diff, ln, latex, Eq, solve, exp,
     simplify
3 # Define symbols
4 N, Np, beta, epsilon = symbols('N Np beta epsilon')
6 # Define the function
7 \ln_t Np = N * \ln(N) - (N - 2*Np) * \ln(N - 2*Np) - Np * \ln(Np)
      - Np * ln(2) - beta * epsilon * Np
9 # Differentiate ln(t_Np) with respect to Np
10 d_ln_t_Np_dNp = diff(ln_t_Np, Np)
# print(latex(d_ln_t_Np_dNp.simplify()))
14 # solve for N_P here: \exp{\left(\beta \epsilon - 1 \right)}
     = \frac{(N - 2 N_p)^2}{2N_p}
final_equation = Eq(exp(beta * epsilon - 1), (N - 2*Np)**2
     (2*Np))
16 Np_solution = solve(final_equation, Np)
print(latex(Np_solution[0].simplify()))
```

We are really interested in the fraction of monomers associated,  $p = \frac{2N_p}{N}$ , so we can divide by N and multiply by 2:

$$p = 1 + \frac{2e^{\beta\epsilon} - \sqrt{(4eN + e^{\beta\epsilon})} 2e^{\beta\epsilon}}{4Ne}$$
 (13)

Not sure where to go from here.

(e) (6 pts) Let us now consider the limits of the fraction of monomers associated. Consider both the limit at high  $(\beta \to 0)$  and low  $(\beta \to \infty)$  temperatures. In terms of energetic interactions and conformational contributions, which effect dominates in either limit?

### 1.5

We want to consider the term:

$$p = 1 + \frac{\frac{\exp(-\beta\epsilon)}{N} - \sqrt{\left(2 + \frac{\exp(-\beta\epsilon)}{N}\right)^2 - 4}}{2} \tag{14}$$

At high temperatures,  $\beta \to 0$ , we can Taylor expand the exponential term to second order:

$$p = 1 + \frac{\frac{1-\beta\epsilon}{N} - \sqrt{\left(2 + \frac{1-\beta\epsilon}{N}\right)^2 - 4}}{2} \tag{15}$$

The term in the square root is then:

$$\left(2 + \frac{1 - \beta \epsilon}{N}\right)^2 - 4 = 4 + \frac{4(1 - \beta \epsilon)}{N} + \frac{(1 - \beta \epsilon)^2}{N^2} - 4 = \frac{4(1 - \beta \epsilon)}{N} + \frac{(1 - \beta \epsilon)^2}{N^2} \tag{16}$$

The former term only scales like  $\sqrt{\frac{1}{N}}$ , so negative term associated with the square root is much larger than the positive term outside. So, I would say that in this high temperature limit, the conformational entropy dominates and the fraction of monomers associated becomes close to 0.

At low temperatures,  $\beta \to \infty$ , the exponential terms in the numerator become negligible, and we are left with:

$$p = 1 + \frac{-\sqrt{(2)^2 - 4}}{2} = 1 + \frac{-\sqrt{0}}{2} = 1 \tag{17}$$

In this low temperature limit, the energetic interactions dominate and the fraction of monomers associated becomes close to 1.

## 2 A diluted magnetic fluid ( 30pts )

Consider a lattice with N lattice sites and n spins in an external magnetic field H that is pointing up. The lattice need not have every site filled with a spin.

- Spins can be either up or down, corresponding to  $s_i = +1$  and  $s_i = -1$ . Spins aligned parallel to the field have an energy -H whereas spins antiparallel have energy H.
- Vacant sites have no spin and thus no interaction with a magnetic field.

You can think of this as a crude model for a paramagnetic fluid that has some non-magnetic impurities. The lattice is connected to a reservoir at temperature T and chemical potential  $\mu$  such that energy E and number of spins n are allowed to fluctuate. We can call this a constrained grand canonical ensemble because the particle number (n) can fluctuate, but cannot exceed N since the lattice has a fixed number of sites. In this case N acts like V because  $V = Nv_{\rm site}$  where  $v_{\rm site}$  is just the volume of a single lattice site.

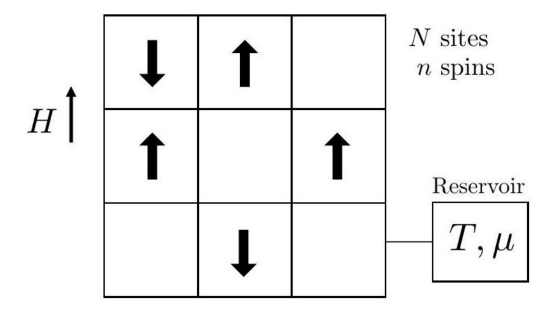


Figure 2: System schematic for the diluted paramagnetic fluid. The example here is showing the case for N=9 and n=5. Note that n fluctuates by exchanging with the reservoir.

(a) (2 pts) What are the definitions of microstate and macrostate for this system?

### 2.1

The microstate is a specific configuration of the lattice, where each site is either filled with a spin or is vacant. The macrostate is a collection of microstates that have the same energy and number of spins.

(b) (8 pts) Derive the closed-form grand canonical partition function  $\Xi(N, T, \mu, H)$ .

Start from either equality below

$$\Xi(N, T, \mu, H) = \sum_{\nu} e^{-\beta E_{\nu} + \beta \mu n_{\nu}} = \sum_{n} Q(N, T, n, H) e^{\beta \mu n}$$

and show that

$$\Xi(N, T, \mu, H) = \left[1 + 2e^{\beta\mu}\cosh(\beta H)\right]^{N}$$

There are multiple ways to do this!

### 2.2

At each site there are 3 possibilities: the site could be vincent, giving an energy of 0, the spin could be aligned with the field, giving an energy of -H, or the spin could be antiparallel to the field, giving an energy of H. The partition function for a single site is then:

$$Q_{\text{site}} = 1 + e^{\beta\mu}e^{-\beta H} + e^{\beta\mu}e^{\beta H} \tag{18}$$

We can factor out the  $e^{\beta\mu}$  from the last two terms and multiply them by  $\frac{2}{2}$ :

$$Q_{\text{site}} = 1 + 2e^{\beta\mu} \left( \frac{e^{-\beta H} + e^{\beta H}}{2} \right) \tag{19}$$

We can recognize the term in the parentheses as the hyperbolic cosine:

$$Q_{\text{site}} = 1 + 2e^{\beta\mu} \cosh(\beta H) \tag{20}$$

This is a partition function for a single site, so for all N sites, we get:

$$\Xi(N, T, \mu, H) = \left[1 + 2e^{\beta\mu} \cosh(\beta H)\right]^{N} \tag{21}$$

(c) (6 pts) Solve for the average fraction of occupied sites in the system  $\langle x \rangle = \langle n \rangle / N$  using an appropriate derivative. What happens as  $H \to \infty$ ? Explain the physical intuition of the limiting behavior.

### 2.3

The relevant thermodynamic potential is the grand free energy,  $W = -kT \ln(\Xi)$ . in its deferential form, this is given by:

$$dW = -SdT - PdV - Nd\mu - MdH \tag{22}$$

So, if we want to solve for the average value of n, we can take the negative derivative of W with respect to  $\mu$ , or equivalently, the derivative of  $\ln(\Xi)$  with respect to  $\beta\mu$ . We get:

$$\frac{\partial \ln(\Xi)}{\partial \mu} = \frac{N}{1 + 2e^{\beta\mu}\cosh(\beta H)} \times 2\beta e^{\beta\mu}\cosh(\beta H) \tag{23}$$

Dividing by N and  $\beta$ :

$$\langle x \rangle = \left( \frac{2e^{\beta\mu} \cosh(\beta H)}{1 + 2e^{\beta\mu} \cosh(\beta H)} \right) \tag{24}$$

The limit of  $\cosh(x)$  as  $x \to \infty$  is  $\frac{e^x}{2}$ , or  $\infty$ . So, as  $H \to \infty$ , the fraction of occupied sites approaches 1. This makes sense, as many sites will want to have spins that align with the field to minimize their energy.

(d) (3 pts) In the reservoir, we have H = 0. If we also know that the fraction of sites occupied in the reservoir is  $x_R$ , then what is  $e^{\beta\mu}$ ?

### 2.4

If in the reservoir, H = 0, then  $\cosh(\beta H) = 1$ . So the fraction of occupied sites in the reservoir is given by:

$$x_R = \frac{2\beta e^{\beta\mu}}{1 + 2e^{\beta\mu}} \tag{25}$$

Solving for  $e^{\beta\mu}$ :

$$e^{\beta\mu} = \frac{x_R}{2\left(\beta - x_R\right)} \tag{26}$$

```
from sympy import *

# divine the symbols of the equation x_R = \frac{2\beta e^{\\ beta\mu}}{1+2e^{\\beta\mu}}

x_R, beta, mu = symbols('x_R beta mu')

eq = Eq(x_R, 2*beta*exp(beta*mu)/(1+2*exp(beta*mu)))

# solve for e^{\\beta\mu}

solution = solve(eq, exp(beta*mu))

print(latex(solution[0].simplify()))
```

(e) (4 pts) Calculate the average magnetisation  $\langle M \rangle$ .

Note the differential form of the grand free energy including the magnetisation

$$dW = -SdT - Pv_{\text{site}} dN - nd\mu - MdH$$

### 2.5

The average magnetisation is given by the derivative of the grand free energy with respect to H:

$$\langle M \rangle = -\frac{\partial W}{\partial H} = \frac{1}{\beta} \frac{\partial \ln(\Xi)}{\partial H}$$
 (27)

This derivative is given by:

$$\frac{\partial \ln(\Xi)}{\partial H} = \frac{2N\beta e^{\beta\mu}\sinh(H\beta)}{2e^{\beta\mu}\cosh(H\beta) + 1} \tag{28}$$

Dividing by  $\beta$ :

$$\langle M \rangle = \frac{2Ne^{\beta\mu}\sinh(H\beta)}{2e^{\beta\mu}\cosh(H\beta) + 1} \tag{29}$$

(f) (3 pts) Calculate the magnetic susceptibility,  $\chi = (\partial \langle M \rangle / \partial H)_{T,N,\mu}$ .

### 2.6

The magnetic susceptibility is given by the derivative of the average magnetisation with respect to H with T, N, and  $\mu$  held constant:

$$\chi = \frac{\partial \langle M \rangle}{\partial H} = \frac{2N\beta \left(2e^{\beta\mu} + \cosh(H\beta)\right) e^{\beta\mu}}{\left(2e^{\beta\mu}\cosh(H\beta) + 1\right)^2}$$
(30)

```
from sympy import *

# Define the symbols for \langle M \rangle = \frac{2N e^{\\
beta \mu} \sinh{\left(H \beta \right)}}{2 e^{\\beta \mu} \\
cosh{\left(H \beta \right)} + 1}

N, beta, mu, H = symbols('N beta mu H')

M = 2*N*exp(beta*mu)*sinh(H*beta) / (2*exp(beta*mu)*cosh(H*beta) + 1)
```

```
# take the deviate with respect to H
6 dM_dH = diff(M, H)
7 print(latex(dM_dH.simplify()))
```

(g) (4 pts) Relate the susceptibility  $\chi$  to the fluctuations in magnetisation  $\sigma_M^2$ . How does  $\sigma_M/\langle M \rangle$  scale with N? This should require little additional work!

### 2.7

Due to the fluctuation dissipation theorem, we can relate the extensive magnetization fluctuations to the susceptibility:

$$\sigma_M^2 = \frac{\chi}{\beta} \to \sigma_M = \sqrt{\frac{\chi}{\beta}} \tag{31}$$

So:

$$\frac{\sigma_M}{\langle M \rangle} = \sqrt{\frac{\chi}{\beta}} \frac{1}{\langle M \rangle} = \sqrt{\frac{\chi}{\beta}} \frac{1}{\frac{2Ne^{\beta\mu}\sinh(H\beta)}{2e^{\beta\mu}\cosh(H\beta) + 1}} = \sqrt{\frac{\chi}{\beta}} \frac{2e^{\beta\mu}\cosh(H\beta) + 1}{2Ne^{\beta\mu}\sinh(H\beta)}$$
(32)

$$\sqrt{\frac{\chi}{\beta}} \tag{33}$$

is an extensive quantity, scaling like  $\sqrt{N}$ , so the ratio  $\sigma_M/\langle M \rangle$  scales like  $1/\sqrt{N}$ .

# 3 Simple Elastic Molecule under Fixed Tension (40 pts)

A crude way to treat a rubber band is by treating it as a chain of linked segments that can take configurations of different lengths. In general we could also assume that the different configurations have different energies.

Assume that there are N total distinguishable segments which each have access to two different conformations: type  $\mathbf{A}$  and type  $\mathbf{B}$ . The types are distinguished by their length and their energy.

- Type A has length a and energy  $\epsilon_a$ .
- Type B has length b and energy  $\epsilon_b$ .

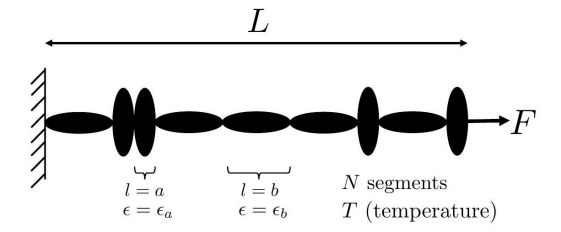


Figure 3: Schematic for the elastic molecule. Note that in this case L can fluctuate by changing segment conformations (not by changing N). The specific example shown is for N=9 and L=4a+5b.

(a) (8 pts) The molecule is placed in a bath at temperature T and extended with a fixed force F. Starting from the Euler equation for the relevant free energy at fixed (N, F, T), use the variational method to derive the probability distribution  $P_{\nu}(N, F, T)$  and show that the partition function Z(N, F, T) in this ensemble is given by:

$$Z(N, F, T) = \sum_{\nu} e^{\beta F L_{\nu}} e^{-\beta E_{\nu}}$$

Note that the Euler equation and fundamental equation for the internal energy are, respectively:

$$E = TS + FL + \mu N$$
$$dE = TdS + FdL + \mu dN$$

### 3.1

We are dealing with a system of a fixed number of segments, N, so particle exchange will not be of concern in our Boltzmann factor. The chain will seek to minimize the energy of a microstate, which adds a factor of  $e^{-\beta E_{\nu}}$ . However, it is also subject to a force, F, which when stretching the chain, will add a factor of  $e^{\beta FL_{\nu}}$  since the chain has more free energy (e.g. how a rubber

band has more potential energy when stretched). So, the total Boltzmann factor for a microstate is given by:

$$e^{\beta F L_{\nu}} e^{-\beta E_{\nu}} \tag{34}$$

The partition function is then given by the sum of the Boltzmann factors for all microstates:

$$Z(N, F, T) = \sum_{\nu} e^{\beta F L_{\nu}} e^{-\beta E_{\nu}}$$
(35)

and the probability for being in a certain state  $P_{\nu}(N, F, T)$  is given by the Boltzmann factor for that state, divided by the partition function:

$$P_{\nu}(N, F, T) = \frac{e^{\beta F L_{\nu}} e^{-\beta E_{\nu}}}{Z(N, F, T)}$$
 (36)

(b) (6 pts) Using the answer to part (a), show that the partition function in the (N, F, T) ensemble is given by

$$Z(N, F, T) = \left[ e^{\beta Fa} e^{-\beta \epsilon_a} + e^{\beta Fb} e^{-\beta \epsilon_b} \right]^N$$

### 3.2

First we will consider the partition function for a single segment:

$$Z_1 = e^{\beta Fa} e^{-\beta \epsilon_a} + e^{\beta Fb} e^{-\beta \epsilon_b} \tag{37}$$

That is, the segment can either be of type A or type B and the partition function is the sum of the Boltzmann factors for each type. The partition function for N segments is then given by:

$$Z(N, F, T) = Z_1^N = \left[ e^{\beta F a} e^{-\beta \epsilon_a} + e^{\beta F b} e^{-\beta \epsilon_b} \right]^N$$
(38)

This can be done because the segments are distinguishable, so the partition function for N segments is just the partition function for a single segment raised to the power of N.

We will now look at a specific case, that yields some very interesting results. Let's take the simplest example, where  $\epsilon_a = \epsilon_b = 0$  and a = 0. Thus, each segment is either vertical (and has no length) or is horizontal (and has length b). In this case, the partition function simplifies to,

$$Z(N, F, T) = \left[1 + e^{\beta F b}\right]^{N}$$

(c) (4 pts) Calculate the extension-force equation of state,  $\langle L \rangle = L(N, F, T)$ .

$$\langle L \rangle = Nb \frac{e^{\beta Fb}}{1 + e^{\beta Fb}}$$

### 3.3

To find the average extension L, we consider that it is given by:

$$\langle L \rangle = N * b * P(B) \tag{39}$$

This can be explained as follows: we are concerned with total number of segments, N. Since we know that a = 0, only segments of type B will contribute to the extension. That's why we multiply by the length of segment B, b and then the probability of a segment being of type B, P(B). The probability of a segment being of type B is given by:

$$P(B) = \frac{e^{\beta Fb}}{1 + e^{\beta Fb}} \tag{40}$$

The denominator is the partition function for a single segment, and the numerator is the Boltzmann factor for a segment being of type B. So, the average extension is given by:

$$\langle L \rangle = Nb \frac{e^{\beta Fb}}{1 + e^{\beta Fb}} \tag{41}$$

(d) (4 pts) Find the equilibrium length  $(L_0)$  of the molecule at zero tension (i.e. at F = 0). Does this answer make sense? Why or why not?

### 3.4

At zero tension, F = 0, the average extension is given by:

$$\langle L \rangle = Nb \frac{e^{\beta Fb}}{1 + e^{\beta Fb}} = Nb \frac{e^0}{1 + e^0} = Nb \frac{1}{2}$$
 (42)

This answer makes sense come because without the force to favor one conformation over the other, the molecule will have an equal probability of being

in either conformation with it's length just given then by the average of the two lengths.

(e) (6 pts) Now we will derive Hooke's law and calculate the spring constant using statistical mechanics!

### 3.5

i. Invert L(N, F, T) to get the force-extension equation of state,  $F(\langle L \rangle, N, T)$ .

### 3.5.1

SymPy gives:

$$F = \left\lceil \frac{\log\left(\frac{L_{avg}}{-L_{avg} + Nb}\right)}{b\beta} \right\rceil \tag{43}$$

```
from sympy import symbols, solve, Eq, exp, ln, latex

# Define symbols
L_avg, N, b, beta, F = symbols('L_avg N b beta F')

# Given equation for L_avg
L_avg_eq = Eq(L_avg, N * b * exp(beta * F * b) / (1 + exp(beta * F * b)))

# Solve the equation for exp(beta * F * b)
exp_beta_Fb_solution = solve(L_avg_eq, exp(beta * F * b))[0]

# Express the solution in terms of F
F_solution = solve(Eq(exp_beta_Fb_solution, exp(beta * F * b)), F)

print(latex(F_solution))
```

ii. Taylor expand  $F(\langle L \rangle, N, T)$  out to first order around the equilibrium chain length from 3d. Physically, when does this approximation apply?

#### 3.5.2

The Taylor expansion of  $F(\langle L \rangle, N, T)$  around the equilibrium chain length is given by:

$$F(\langle L \rangle, N, T) \approx F(L_0, N, T) + \frac{\partial F}{\partial L}\Big|_{L=L_0} (\langle L \rangle - L_0) + \dots$$
 (44)

We know that first term is zero, because the force is zero at the equilibrium length. We are interested in evaluating the derivative of F with respect to L at  $L = L_0$ . Rewriting our expression for F:

$$F = \frac{1}{b\beta} \left( \ln(\langle L \rangle) - \ln(Nb - \langle L \rangle) \right) \tag{45}$$

Taking the derivative of F with respect to L:

$$\frac{\partial F}{\partial L} = \frac{1}{b\beta} \left( \frac{1}{\langle L \rangle} + \frac{1}{Nb - \langle L \rangle} \right) \tag{46}$$

Evaluating this at  $L = L_0$ , we get:

$$\left. \frac{\partial F}{\partial L} \right|_{L=L_0=Nb\frac{1}{2}} = \frac{4}{Nb^2\beta} \tag{47}$$

```
from sympy import diff, symbols, log, solve, Eq, latex,
    simplify

# Define symbols
L_avg, L_O, N, b, beta, F = symbols('L_avg L_O N b beta F')

# Given F(L_avg) equation from previous solution
F_L = log(L_avg / (-L_avg + N * b)) / (b * beta)

# Calculate the first derivative of F with respect to L_avg

# Calculate this derivative at the inital length ofL_O = N b \
    frac{1}{2}

dF_dL_at_LO = dF_dL.subs(L_avg, N * b / 2)

print(latex(dF_dL_at_LO.simplify()))
```

iii. Extract the spring constant  $K_s$  by inspection.

### 3.6

The spring constant is given by the derivative of the force with respect to the extension at the equilibrium length. So, it is what we just calculated:

$$K_s = \frac{4}{Nb^2\beta} \tag{48}$$

(f) (4 pts) Relate the fluctuations in length  $\sigma_L^2$  at F=0 to the spring constant,  $K_s$ . Explain why this relationship physically makes sense.

### 3.7

We can relate the fluctuations in length to the spring constant using the fluctuation dissipation theorem:

$$\sigma_L^2 = \frac{kT}{K_s} \tag{49}$$

This relationship makes sense because the spring constant is a measure of how stiff the chain is, and the fluctuations in length are a measure of how much the chain can move around. So, the stiffer the chain, the less it will fluctuate.

(g) (8 pts) For this more simplified model of the chain where each segment is either vertical or horizontal, we can analyze the behavior using a fixed energy and length ensemble (i.e. the microcanonical ensemble). The partition function is given by  $\Omega(N, L, E)$ .

### 3.8

i. Calculate the microcanonical partition function at length L=nb, where n is the number of horizontal segments.

### 3.8.1

We want to calculate the number of ways to arrange n horizontal segments (which will have a length b, but doesn't matter here) in a chain of N segments. This is given by the binomial coefficient:

$$\Omega(N, L, E) = \binom{N}{n} = \frac{N!}{n!(N-n)!}$$
(50)

ii. Show that your answer to **3d** maximizes the entropy.

### 3.8.2

The entropy is given by:

$$S = k \ln(\Omega) \tag{51}$$

We want to evaluate  $ln(\Omega)$ :

$$\ln(\Omega) = \ln\left(\frac{N!}{n!(N-n)!}\right) = \ln(N!) - \ln(n!) - \ln((N-n)!)$$
 (52)

We can use Stirling's approximation to simplify this:

$$\ln(\Omega) \approx N \ln(N) - n \ln(n) - (N - n) \ln(N - n) \tag{53}$$

The entropy is then given by:

$$S = k \ln(\Omega) \approx k \left( N \ln(N) - n \ln(n) - (N - n) \ln(N - n) \right) \tag{54}$$

We can now differentiate this with respect to n to find the most-probable value of n.

$$\frac{\partial S}{\partial n} = k \left( -\log\left(n\right) + \log\left(N - n\right) \right) \tag{55}$$

Setting this equal to zero:

$$0 = -\log(n) + \log(N - n) \tag{56}$$

Bringing one of the terms to the other side:

$$\log\left(n\right) = \log\left(N - n\right) \tag{57}$$

Exponentiating both sides:

$$n = N - n \tag{58}$$

Solving for n:

$$n = \frac{N}{2} \tag{59}$$

or we have  $n = \frac{L}{b}$ . So, we plug in and solve for L:

$$L = nb = \frac{N}{2}b\tag{60}$$

which is the same as the equilibrium length we found in 3d. So, the answer to 3d maximizes the entropy.

iii. Verify that you obtain the same force-extension equation of state F(N, L, T) in this ensemble.

### 3.8.3

As showed earlier, this ensemble has the equilibrium length with the total force F being zero. The force-extension equation of state is given by:

$$F = \frac{\partial E}{\partial L} = \frac{1}{b\beta} \left( \ln(\langle L \rangle) - \ln(Nb - \langle L \rangle) \right) \tag{61}$$

But when we set the average length to the equilibrium length of  $L_0 = \frac{N}{2}b$ , we get:

$$F = \frac{1}{b\beta} \left( \ln \left( \frac{N}{2} b \right) - \ln \left( Nb - \frac{N}{2} b \right) \right) = 0 \tag{62}$$