We develop a statistical model for a confined homopolymeric chain molecule based on a monomer grand ensemble representation. The molecule is subject to a confining external field, a backbone interaction, and an attractive interaction between any pair of monomers. An exact minimum principle for the thermodynamics of the backbone in an external field is obtained, and a controlled mean field approximation results in a modified minimum principle from which relevant physical quantities such as monomer density can be found. We explore the limit in which the chain is subject to tight confinement, and make a preliminary investigation of a prototypical system.

Keywords: confined polymers; folding; long-range interactions

1. Introduction
The question of how linear macromolecules like proteins fold inside a cavity is of great biological relevance. Indeed it is now recognised that large proteins are not capable of correctly fold in the crowded cell’s environment, where the volume fraction occupied by macromolecules can be of the order of 20–30% [1], but they need some kind of cavities to fold, where they are isolated from the rest of the cell’s environment, and can avoid interactions with other molecules which would cause misfolding and aggregation [2]. Moreover, in vitro experiments where the effects of confinement on the thermodynamics of protein folding were directly monitored, seem to indicate that confinement increases the stability of the native states [3].

The simplest representation of biopolymers under confinement, is one where the molecule is modelled by a classical chain under an external field. The work presented in this paper aims at the development of a consistent and well grounded perturbative approach to the theory of confined polymer folding. Studies of polymers have of course been extensive, and useful relationships between polymer melts and fluids established [4], as well as between single polymers and fluids. This paper represents an initial investigation, aimed at understanding which concepts borrowed from bulk fluid studies [5,6] still remain relevant in a polymer setting. The classical theory of fluids in thermal equilibrium is a highly developed discipline. Along with specific physically motivated approximations have come tools of more general validity and utility.

Our focus will be on a single polymer chain confined by external forces, and to minimise the needed information input, on single homopolymers. We will also attend in the main to idealised models in which the polymer is simply a chain of unit monomers of a few degrees of freedom, but will indicate how this restriction can be rewardingly removed, on the way to a realistic polymer representation.

We will aim at both analytic simplicity and reasonable suitability for the ultimately necessary computational procedures. For the former, we will work in a ‘monomer grand-ensemble’ [7,8] in which the number of monomers per polymer is distributed, but will show that this need not be a drawback. For the latter, we will favour minimum principles to be able to better control computations. This does restrict the category of systems to be studied e.g. to purely attractive pair interactions, and will be generalised in later work, soon to be reported, as such restrictions are removed.

2. The reference system
The statistical mechanics of a free classical chain molecule, interpreted as a random walk is an old, old topic, although less attention has been paid to the
effect of a structured external field, in part because the formal simplification afforded by translation invariance is then not available. In this case, the next neighbour pair energy then depends separately on the locations of each of the monomers in the pair, rather than on just their separation (in the point monomer model). We will tackle this problem by first (Section 2.1) concealing the discreteness of the monomer population number, and then (Section 2.2) concealing the inherent required inversion of the pair Boltzmann factor kernel.

2.1. Notation

Let us be a bit more explicit. We have in mind an ordered chain of \( N \) equivalent monomers, the \( j \)th being specified by its degrees of freedom \( r_j \). The order is maintained by a symmetric next neighbour interaction potential of Boltzmann factor \( w(r_j, r_{j+1}) \), depending on course on the inverse temperature \( \beta \). Any two monomers can also interact via an interaction Boltzmann factor \( e(r_i, r_j) = \exp[-\beta\phi(r_i, r_j)] \) and, crucially for the applications we have in mind, the polymer is constrained by an external potential \( u(r_j) \).

It is now convenient to imagine that the homopolymer in question is both in thermal equilibrium and in number equilibrium, i.e. that it is the result of monomer addition and absorption from a bath of non-interacting monomers (Figure 1). The reaction equilibrium is analogous to that of the grand canonical ensemble for a fluid, but with a significant difference. Suppose that \( Q^m \) is the monomer canonical partition function in its centre-of-mass coordinate system, \( Q^p_\text{N} \) of the \( N \)-monomer polymer (\( N \geq 1 \), to recognise an object as a polymer). Then if the full system contains \( N' \) monomers, \( N \) of which are bound together, in a volume \( V \), the full system partition function will be:

\[
Q^\text{tot}_N = \sum_{N \geq 1} \frac{(VQ^m)^{N'-N}}{(N'-N)!} Q^p_\text{N}.
\]

Since the monomers in the absence of a polymer would have a partition function:

\[
Q^p_\text{N} = \frac{(VQ^m)^{N}}{N!},
\]

that attributed to the polymer will take on the form:

\[
\Xi^{(p)}[\zeta_0] = \lim_{N \to \infty} \frac{N!}{(VQ^m)^N} Q^\text{tot}_N = \sum_{N \geq 1} \zeta_0^N Q^p_\text{N}, \quad \text{where } \zeta_0 = (N'/V)/Q^m.
\]

The obvious analogy with a fluid grand partition function (with the weight \( 1/N! \) excised) will be very useful indeed.

The computation of \( \Xi^{(p)}[\zeta_0] \), which generates all thermodynamics and expectations in a thermal ensemble, is of course too general to be explicitly solvable, except in very special circumstances. Let us therefore start with the evaluation of \( \Xi^{(p)}[\zeta_0] \) for what may be regarded as the backbone of the polymer, that in which the arbitrary pair mutual interaction \( \phi(r_j, r_i) \) is set equal to zero. In this case, we have at once:

\[
\zeta_0^N Q^p_\text{N} = \int \prod_{i=1}^{N-1} w(r_i, r_{i+1}) \xi(r_1) \xi(r_2) \ldots \xi(r_{N-1})
\]

\[
\times w(r_{N-1}, r_N) \xi(r_N) \, dr^N
\]

where

\[
\xi(r) = \zeta_0 \exp[-\beta u(r)] = \exp[\beta \mu(r)]
\]

or regarding \( w(r, r') \) as the kernel of an integral operator \( w \), \( \xi(r) \delta(r-r') \) as that of the diagonal operator \( \xi \), and the symbol \( |1\rangle \) denoting the vector whose components are 1,

\[
\Xi^{(p)}[\xi] = \langle 1 | (\xi^{-1} - w)^{-1} | 1 \rangle,
\]

subject of course to convergence of the series (4). The corresponding ‘grand potential’ is:

\[
\Omega^{(p)}[\xi] = -\frac{1}{\beta} \ln(\langle 1 | (\xi^{-1} - w)^{-1} | 1 \rangle)
\]
and as an immediate consequence the monomer density is given by

\[ n(r) = -\beta \xi(r) \delta \frac{\delta}{\delta \xi(r)} \Omega^{(\phi)} [\xi] \]  
\[ = \langle 1 | (\xi^{-1} - w)^{-1} | r \rangle \langle r | (\xi^{-1} - w)^{-1} | 1 \rangle \xi^{-1}(r) \]
\[ / \langle 1 | (\xi^{-1} - w)^{-1} | 1 \rangle. \]  

(9)

The absence of the statistical weight \(1/N!\) in (4) suggests an unusually broad distribution of monomer number \(N\) about its mean, in the ensemble. In Appendix 1, we examine the form of this distribution, its consequences, and the nature of required correction terms.

2.2. Minimum principle

Expression (7) for the backbone grand potential is very concise, but it requires the inversion of an integral operator, \((\xi^{-1} - w)\), and so is technically non-trivial, although computationally not a great impediment. More importantly, it makes unfeasible the direct use of (7) as a potential building block for the full interacting system. Fortunately, (7) can be constructed, making use of a minimum principle in which no such inverses appear. Doing so depends upon the fact that if \(K\) is positive semidefinite, and \(a\) and \(\psi\) are arbitrary, then according to the Schwartz inequality we have

\[ \langle a | K^{-1} | a \rangle \langle \psi | K | \psi \rangle = \langle K^{-1/2} a | K^{-1/2} a \rangle \langle K^{1/2} \psi | K^{1/2} \psi \rangle \]
\[ \geq \langle K^{-1/2} a | K^{1/2} \psi \rangle^2 = \langle a | \psi \rangle^2. \]  

(11)

so that \(\langle a | K^{-1} | a \rangle \geq \langle a | \psi \rangle^2 / \langle \psi | K | \psi \rangle\). Then indeed

\[ \max_{\psi} \frac{\langle a | \psi \rangle^2}{\langle \psi | K | \psi \rangle} = \langle a | K^{-1} | a \rangle \text{ at } K \psi = \frac{\langle \psi | K | \psi \rangle}{\langle a | \psi \rangle} a, \]  

(12)

\[ \beta \Omega^{(\phi)} [\xi] = \min_{\psi} \ln \langle \psi | \xi^{-1} - w | \psi \rangle - 2 \ln \langle 1 | \psi \rangle. \]  

(13)

The representation (13), it must be emphasised, is exact. There is no explicit inverse contained therein, and this indeed allows it to be used as a first step in analysis of the backbone with non-neighbour pair forces.

3. The mean field strategy

3.1. Exact results

Having the backbone of the polymer in an external field potentially under control, we now approach our real objective: to find the effect of mutual interactions, \(\phi_1(r, r')\), between any two monomers at \(r\) and \(r'\).

The most common leading estimate in such situations, termed mean field, is one in which the mean value of the sum of pair interactions acting on a given unit is taken as an additional external field. This acts self-consistently on each unit, and so control of the non-interacting system then suffices. For classical fluids, Widom’s insertion theorem [9] tells us that if the calculation is done instead for each member of the ensemble – fluctuations hence automatically included – and then averaged over the ensemble, it becomes exact in the sense that

\[ n(r) = \left\{ \exp \left[ \beta (\mu(r) - \int \phi_1(r, r') \hat{n}(r') \, dr' \right) \right\}, \]  

(14)

where

\[ \hat{n}(r) = \sum_i \delta(r - r_i) \]  

(15)

is the microscopic, per configuration, particle density. Mean field thus amounts to replacing \(\hat{n}(r')\) by \(n(r') = n(r')\). Carrying out corrections however leads to questions about the pair distribution, which can get quite involved. For our purposes, another exact representation is preferable, in which the interaction is replaced by a suitable average over an equivalent fluctuating external field. This is of very general validity, but it applies rigorously only to the additions of purely attractive (negative definite) pair forces, which certainly can distort local properties of a real system. In the case of a polymer backbone, which already includes next neighbour pair forces, this assumption is not lethal, and we can anticipate a reasonable representation of global structure. Suppose then that, quite generally, we set

\[ \phi_1(r, r') = -\phi(r, r') \]  

(16)

for convenience (\(\phi\) is now positive definite as a continuous matrix) and imagine that \(\phi\) is added to a reference system all of whose properties are known.

\[ \Xi[\mu, \phi] = \sum_n \int \ldots \int W_0^{(N)}(r_N, \mu) \exp \left[ -\frac{\beta}{2} \sum_{i,j} \phi(r_i, r_j) \right] \, dr_N \]  
\[ = \sum_n \int \ldots \int W_0^{(N)}(r_N, \mu) \exp \left[ -\frac{\beta}{2} \sum_i \phi_D(r_i) \right] \times \exp \left[ \frac{\beta}{2} \sum_{i,j} \phi(r_i, r_j) \right] \, dr_N \]  
\[ = \sum_n \int \ldots \int W_0^{(N)}(r_N, \mu) \exp \left[ -\frac{\beta}{2} \sum_i \phi_D(r_i) \right] \times \exp \left[ \frac{\beta}{2} \int \hat{n}(r') \phi(r, r') \hat{n}(r') \, dr' \right] \, dr_N \]  

(18)

(19)

where \(\sum\) omits the \(i = j\) contribution, \(\phi_D\) is the diagonal part of the matrix \(\phi(r_i, r_j)\).
The device of Kac, Siegert, Hubbard and Stratonovich [10–13] is to represent the Gaussian in (17) (in obvious notation) as a functional Laplace transform

$$\exp\left(\frac{\beta}{2} \vec{n} \cdot \vec{\phi} \right)$$

$$= \int \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right)$$

$$\times \exp(-\beta \vec{v} \cdot \vec{n}) D\vec{v} / \int \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}$$

$$= \int \exp\left[-\beta \sum_n \vec{v}(r_n)\right] \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}$$

$/\int \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}.$

(20)

Since $\sum_n \int \ldots \int W_r^{N}(v^N, \mu) d^N = Z[\mu]$, Equation (17) can thereby be rewritten as

$$\Xi[\mu, \phi] = \int Z[\mu - \frac{1}{2} \phi_D - v] \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}$$

$$/\int \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}.$$

(22)

with a possible interpretation that the interaction $-\phi$ has been replaced by an ensemble average over a fluctuating external field $\vec{v}(r)$, serving as a sort of graviton shuttling back and forth between units, and in fact, particle configuration space has now been transformed to field space. The physical interpretation is comfortable, but more importantly, it leads to a new method of evaluation as well, and applies to fluids, polymers, . . .

The kernel of (22) is a Boltzmann factor in field space, and so we may define a field average as

$$\langle(G[v])\rangle = \int \mathcal{G}[v] \Xi[\mu - \frac{1}{2} \phi_D - v] \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}$$

$/\Xi[\mu, \phi].$

(23)

A very suggestive consequence of this notation follows from the observation that for the density $n(r)$,

$$n(r) = \frac{1}{\Xi[\mu, \phi]} \delta \mu(r) \Xi[\mu, \phi]$$

$$= -\frac{1}{\delta \mu(r)} \Xi[\mu - \frac{1}{2} \phi_D - v] \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}$$

$/\Xi[\mu, \phi] \int \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}$

(24)

or integrating by parts in v-space (assuming the absence of boundary terms)

$$n(r) = -\int \phi^{-1}(r) \Xi[\mu - \frac{1}{2} \phi_D - v] \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}$$

$/\Xi[\mu, \phi] \int \exp\left(-\frac{\beta}{2} \vec{v} \cdot \vec{\phi}^{-1} \vec{v} \right) D\vec{v}$

(26)

identifying $-\phi n(r)$ as the ‘mean field’ $\langle \phi(r) \rangle$. Equation (27) is exact, but depends very much on the details of the system under study e.g. in (26). Exact calculations in function space are out of the question, and so it is the approximate evaluation of $\langle \phi(r) \rangle$ that we must attend to.

3.2. A mean field approximation

There are any number of ways of setting up approximation sequences for the evaluation of $\langle \phi(r) \rangle$. We will adopt one which is fairly obvious in the context of (22) and (26), with the great advantage of a very pictorial leading order – still appearing as a minimum principle – and a physically transparent first order. It goes as follows. If the field Boltzmann factor of (23) is sharply peaked about a function $\tilde{\phi}(r)$, then of course (27) becomes simply

$$n(r) = -\phi^{-1} \tilde{\phi}(r),$$

(28)

and the field $\tilde{\phi}$ must now satisfy (\Omega is again the grand potential $-\beta^{-1} \ln \Xi$)

$$\frac{\delta}{\delta \tilde{\phi}(r)} \left( \Omega_0 \left[ \mu - \frac{1}{2} \phi_D - \tilde{\phi} \right] \right) + \frac{1}{2} \tilde{\phi} \cdot \vec{v} \cdot \phi^{-1} \vec{v} = 0,$$

(29)

or simply

$$n_0 \left( r | \mu - \frac{1}{2} \phi_D - \tilde{\phi} \right) = -\phi^{-1} \tilde{\phi}(r),$$

(30)

the unsurprising result (comparing with (28)) that $n(r)$ is equal to the ‘bare’ density $n_0$ in the presence of $\tilde{\phi}$, to within an additional shift of $\frac{1}{2} \phi_D$. But in addition, we now have in leading order approximation the very explicit

$$\Omega_1(\mu, \phi) = \Omega_0 \left[ \mu - \frac{1}{2} \phi_D - \tilde{\phi} \right] + \frac{1}{2} \tilde{\phi} \cdot \vec{v} \cdot \phi^{-1} \vec{v}$$

(31)

which, since (29) represents the minimisation of the kernel of (24), takes the variational form

$$\Omega_1(\mu, \phi) = \text{Min}_0 \left[ \mu - \frac{1}{2} \phi_D - \tilde{\phi} \right] + \frac{1}{2} \tilde{\phi} \cdot \vec{v} \cdot \phi^{-1} \vec{v}. $$

(32)
Furthermore, consistency is established by noting that by virtue of (32), $\Omega_1$ of (31) implies
\[
 n(r) = -\frac{\delta \Omega_1}{\delta \mu(r)} = -\phi^{-1} \bar{v}(r), \quad (33)
\]
reproducing (28).

To apply (32) to the single homopolymer under discussion, we need only insert (13), serving as $\Omega_0$, into (32). We then have
\[
 \beta \Omega_1[\mu, \phi] = \text{Min}_{n, \phi} \left[ \frac{\beta}{2} v \cdot \phi^{-1} v - 2 \ln(1|\psi) 
 + \ln\langle \psi|\exp[-\beta(\mu - v)] - w|\psi \rangle \right], \quad (34)
\]
where $\mu'(r) = \mu(r) - \frac{1}{2} \phi D(r)$. Equation (34) can be simplified by finding and eliminating the field $\psi$, and replacing $v$ by $-\phi n$. To do so we observe that
\[
 0 = \frac{\delta \Omega_1}{\delta \psi} = \phi^{-1} v + \exp[-\beta(\mu - v)] \psi^2 
/\langle \psi|\exp[-\beta(\mu - v)] - w|\psi \rangle. \quad (35)
\]
Integrating over the implicit $r$, with $N = \int n(r) dr$, then
\[
 N = \frac{\langle \psi|\exp[-\beta(\mu - v)]\rangle|\psi \rangle}{\langle \psi|\exp[-\beta(\mu - v)] - w|\psi \rangle}, \quad (36)
\]
reducing (35) to
\[
 n = N \exp[-\beta(\mu - v)] \psi^2 /\langle \psi|\exp[-\beta(\mu - v)]\rangle. \quad (37)
\]
Since (37) is homogeneous of degree 0 in $\psi$, we are free to adopt the normalisation
\[
 \langle \psi|\exp[-\beta(\mu - v)]\rangle = N \quad (38)
\]
converting (37) to
\[
 n = \exp[-\beta(\mu - v)] \psi^2 \quad (39)
\]
(leading back to (38)), and consequently replace (34) by the simple
\[
 \beta \Omega_1[\mu, \phi] = \text{Min}_n \left[ \frac{\beta}{2} n \cdot \phi n - 2 \ln\left(1|n^{1/2} \exp\left[\frac{\beta}{2}(\mu' + \phi \cdot n)\right]\right) 
+ \ln(n^{1/2}) \left(I - \exp\left[\frac{\beta}{2}(\mu' + \phi \cdot n)\right] w 
\times \exp\left[\frac{\beta}{2}(\mu' + \phi \cdot n)\right]|n^{1/2}\right) \right] \quad (40)
\]
appearing as a $\mu$-dependent density functional, reminiscent of ‘statistical models’ [14–16], of polymers. Equation (40) is our main result, valid in the mean field level. The role of the interaction $-\phi$ is transparent: the external potential is augmented by the mean field: $\mu \to \mu' + \phi n$, and the explicit energetic component $-\frac{1}{2} n \cdot \phi n$ subtracted out. Equation (40) is of course a density functional representation with the density profile dependence on $\mu$ determined by dropping the implicit $\phi$ and making $\mu$ explicit, by
\[
 \delta \Omega_1[\mu, n]/\delta n(r) = 0. \quad (41)
\]
By definition, see Equation (40), we have ignored the fluctuations of the potential field $\bar{v}(r)$ about its mean $\bar{v}$. Corrections must then correspond to including the fluctuations to some model order. This is analogous to including capillary waves in the description of a two-fluid interface, and they do make both energetic and entropic contributions. In Appendix 2, we show briefly how to obtain the effect of potential field fluctuations to Gaussian order, but leave resulting numerical contributions to the future.

4. Numerical implementation

The minimisation required in (40) to obtain numerically the profile $n(r)$ can be carried out in a number of ways. However, in many-monomer systems, which are of major interest, there is an important preliminary observation. In Appendix 3, we analyse the system behaviour as the monomer number increases at fixed external confinement potential. We show that, unsurprisingly considering the floppiness of the chains we are working with at this introductory stage, increasing mean monomer number serves only to increase the monomer density uniformly over the confinement volume. It is therefore only the normalised monomer density $p(r) = n(r)/N$ that is relevant.

In terms of $p$ the minimisation functional (40) becomes
\[
 \Omega_1[\mu, \phi] = \text{Min}_p \left[ \frac{N^2}{2} p \cdot \phi p - 2 \ln\left(1|p^{1/2} \exp\left[\frac{\beta}{2}(\mu' + N\phi p)\right]\right) 
+ \ln(p^{1/2}) \left(I - \exp\left[\frac{\beta}{2}(\mu' + N\phi p)\right] w 
\times \exp\left[\frac{\beta}{2}(\mu' + N\phi p)\right]|p^{1/2}\right) \right] \quad (42)
\]
Once we have determined $p(r)$, the expression to compute the particle number and the true density are
\[
 n(r) = \frac{p(r)}{(p(r)^{1/2})^2} \quad (43)
\]
\[
 N = \frac{p(r)^{1/2}}{(p(r)^{1/2})^2} \quad (44)
\]

For the numerical implementation, we considered a chain confined by hard walls, with harmonic interactions bounding neighbouring monomers, and a long-range step potential active only within a given distance between the monomers. As expected, for the
reference system the probability density converges to a distribution centred in the middle of the box. When the attractive potential $\phi$ is turned on the distribution becomes more peaked (Figure 2).

We also looked at the effect of introducing an asymmetry in the external potential: a uniform, gravitational, potential along one of the axes. As expected the minimisation of the functional converges to a probability density off-centred with respect to $z$ and centred in the middle of the box with respect to $x$ and $y$ (Figure 3).

Even if the theoretical derivation of Equation (40) is valid only for strictly positive definite $\phi$, in the algorithm we can invert the sign of the long range interaction and make it repulsive, instead of just attractive. We can therefore test our method beyond the known validity of the approximations used, taking a repulsive long-range interaction. The method still holds, provided we consider repulsive interactions which are not too strong. As expected a repulsive $\phi$ gives a less peaked density compared to the reference system (Figure 4).

The monomer interactions we consider are both repulsive and attractive. Under this condition we could expect the polymer to undergo a coil-globule transition as the strength of the attractive interaction overcomes the repulsion [17].

In our system, when no long-range interaction is present, the monomers interact harmonically, and the polymer configuration should be that of an extended coil, with the monomers sitting at their equilibrium separations. When the long-range interaction $\phi$ is turned on the monomers start feeling a pure attraction, to all other monomers. There will never be a balance between attraction and repulsion because the repulsion acts only between nearest neighbours, while the attraction is to all other particles in the system. On energetic grounds alone, we would then expect our system to collapse to a globular conformation as soon as the long-range attraction is introduced, even when this is very weak. However, the increased entropy of an open structure softens the rigidity of this statement. Through direct Monte Carlo simulations of a model polymer inside a spherical cavity we indeed verified this behaviour for an attractive $\phi$. When repulsive long-range interactions are introduced instead, the polymer becomes somehow rigid and tends to wrap around in the cavity using all available space to become as close as possible to a straight rod.
With the method developed in this paper, we have systematically looked at the effect of turning on the long-range interaction, performing minimisations at regular increments. In Figure 5 we plot one of the coefficients of the exponent of the gaussian density (we have chosen $B_x$, but results are independent of this choice) as a function of $c_{30}$. We observe a highly non-linear dependence, indicating a rapid compactification of the polymer as $c_{30}$ increases.

![Figure 4](image1.png)

**Figure 4.** Left: interaction between neighbouring particles along the chain: harmonic potential and repulsive long-range interaction. Right: probability profiles at $y = z = L/2$ with (solid) and without (dashed) repulsive core.

![Figure 5](image2.png)

**Figure 5.** The coefficient $B_x$ measured in terms of the strength of the long-range attractive interaction $\phi$. For a Gaussian density like the one we have, a measure of $B_x$ gives an indication of how peaked the density distribution is. We observe a strong non-linear dependence, indicating a rapid compactification of the polymer as $\phi$ increases.

When looking at the details of the polymer configuration, proper quantities to measure are the end-to-end distance or the radius of gyration. At the present stage we measure the average density only. Other quantities are accessible with our formulation, but their theoretical treatment, although straightforward, is lengthy and will involve completely new numerical implementations. This is beyond the scope of this paper, where we want to establish the basic setting for the general formulation of confined polymer behaviour in terms of a minimisation principle and of mean field.

5. Concluding remarks

The simplifying assumptions attend more directly to the physics, and these assumptions depend very much on the nature of the system to be studied. Taking these assumptions in order, we first embedded our system in a monomer grand ensemble. Since preprocessing assures that in practice one does not deal with the resulting extreme polydispersity, a fixed $N$ ensemble is more relevant than fixed $\zeta$. The corresponding inverse mapping has been attended to on numerous occasions (see e.g. [18]). The same formalism is indeed available here (hinted at in Equations (59) and (60)).

Another restriction was to attractive long-range forces. We found however that the profile equation could indeed be pushed into the partially repulsive regime, although the validity of the minimum principle was in question; this is closely related to the functional Fourier transform for the repulsive component, likewise under uncertain control. An alternative approach
lies in the use of the mean spherical model [19,20] and its extensions. This is the aim of ongoing research.

Of course, there is the implicit assumption that pair forces suffice, whereas the action of pairs on singlets is a frequent important observation, leading e.g. to dihedral angle dependence in chains. Typically (see e.g. [21] for a very primitive example) one can simply create a multi-unit monomer to encompass only such forces, which then appear once more as pair forces.

Most bluntly, our approach has been restricted to homopolymers. Since the set of degrees of freedom of a monomer can include monomer type, this is no restriction at all if one is studying the effect on the full population of an assumed relative frequency of occurrence of next neighbours hetero-pairs. However, if we attribute a sense to the chain and the AB frequency differs from the BA frequency, the very conventional symmetry of the operator \( w \) is lost, and with it, the possibility e.g. of a specific long sequence of monomer types. The case of non-symmetric \( w \) has indeed been studied [22], but exercising the kind of control that we have in our current formulation remains a challenge.

References

[1] R.J. Ellis, Curr. Opin. Struct. Biol. 11, 114 (2001).
[2] D. Thirumalai, D.K. Klimov, and G.H. Lorimer, PNAS 100, 11195 (2003).
[3] D.K. Eggers and J.S. Valentine, J. Mol. Biol. 314, 911 (2001).
[4] P.G. de Gennes, Scaling Concepts in Polymer Physics (Cornell University Press, Ithaca, NY, 1979).
[5] I.M. Lifshits, Sov. Phys. JETP 55, 2408 (1968).
[6] S. Pasquali and J.K. Percus, J. Chem. Phys. 125, 064906 (2006).
[7] H.L. Frisch and J.K. Percus, J. Phys. Chem. B 105, 11834 (2001).
[8] J.K. Percus, J. Stat. Phys. 106, 357 (2002).
[9] B. Widom, J. Chem. Phys. 39, 2808 (1963).
[10] M. Kac, in Applied Probability, edited by C.A. MacColl (McGraw Hill, New York, 1957).
[11] A.J.F. Siegert, Physica (suppl.) 26, 530 (1960).
[12] J. Hubbard, Phys. Rev. Lett. 3, 11 (1959).
[13] R.L. Stratonovich, Sov. Phys. Dokl. 2, 416 (1958).
[14] G. Giacomin, Random Polymer Models (World Scientific, Singapore, 2007).
[15] P. Flory, Statistical Mechanics of Chain Molecules (Wiley, New York, 1969).
[16] M. Huggins, J. Phys. Chem. 46, 151 (1942).
[17] D. Poland and H.A. Sheraga, Theory of Helix-Coil Transitions in Biopolymers (Academic Press, New York, 1970).
[18] J.L. Lebowitz, J.K. Percus, and L. Verlet, Phys. Rev. 153, 2506 (1967).
[19] J.K. Percus and G.J. Yevick, Phys. Rev. 136, 290 (1964).
[20] J.L. Lebowitz and J.K. Percus, Phys. Rev. 144, 251 (1966).
[21] K.K. Muller-Nedebock, H.L. Frisch, and J.K. Percus, Phys. Rev. E 67, 011801 (2003).
[22] H.L. Frisch and J.K. Percus, Phys. Rev. E 64, 011805 (2001).
[23] R. Rajaraman, Phys. Rep. C 21, 227 (1975).
[24] J.K. Percus, Physica A 172, 1 (1991).
[25] R. Jentzsch, Crelles J. 141, 235 (1912).

Appendix 1. Number distribution

We first observe from (9) that if \( \hat{N} \) denotes the monomer number in a given configuration, then

\[
N = \langle \hat{N} \rangle = \int n(r) \, dr = (1/(\chi^{-1} - w)^{-1}(\chi^{-1} - w)^{-1}1)/\Xi^{(p)}
\]

\[
= (\chi^{1/2}/(1 - \chi^{1/2}w\chi^{1/2} - 2\chi^{1/2}/\chi^{1/2}/(1 - \chi^{1/2}w\chi^{1/2} - 1)\chi^{1/2})
\]

\[
= 1/(1 - \lambda_0) + O(1/(1 - \lambda_1)),
\]

where \( \lambda_0 = \lambda_{\text{max}}(\chi^{1/2}w\chi^{1/2}) \) and \( \lambda_1 \) is the next largest eigenvalue; \( \lambda_1 < \lambda_0 \) in a confined system. If \( N \) is large, then

\[
\Xi^{(p)}(\xi) = (\chi^{1/2}/(1 - \chi^{1/2}w\chi^{1/2} - 1)\chi^{1/2})
\]

is dominated by the ‘resonance’ at \( \lambda_0 \), so that if

\[
(\chi^{1/2}w\chi^{1/2})\psi_{\lambda_0} = \lambda_0\psi_{\lambda_0}
\]

with normalised \( \psi \), then

\[
\Xi^{(p)}[\xi] \sim \frac{1}{1 - \lambda_0} (\psi_{\lambda_0}\chi^{1/2})^2.
\]

There are now two consequences. On the one hand, we have

\[
\langle \exp(i\theta\hat{N}) \rangle = \exp(i\theta\psi_0)\Xi^{(p)}(\psi_0)/\Xi^{(p)}[\xi]
\]

\[
= \Xi^{(p)}[\xi]\exp(i\theta)/\Xi^{(p)}[\xi]
\]

\[
\cong (1 - \lambda_0)/(1 - \lambda_0\exp(i\theta)),
\]

so that the distribution of \( \hat{N} \) is given by

\[
\beta(N) = \text{cof} [\exp(i\lambda_0\theta)]/(1 - \lambda_0)/(1 - \lambda_0\exp(i\theta)) = \hat{N}_0^\lambda(1 - \lambda_0)
\]

the very broad geometric distribution. It would appear that the \( N \)-ensemble must give a very poor representation of a given \( N \). But on the other hand, we have from (5)

\[
Q_N = c_0^{-N}(\chi^{1/2}/(\chi^{1/2}w\chi^{1/2} - 1)\chi^{1/2})
\]

\[
\cong c_0^{-N}N^{-1}(\psi_{\lambda_0}\chi^{1/2})^2,
\]

so that if \( F_N \) is the canonical Helmholtz free energy,

\[
\beta F_N - \beta \Omega^{(p)} = -\ln Q_N + \ln \Xi^{(p)}[\xi]
\]

\[
\cong N\ln c_0 - (N - 1)\ln \lambda_0 - \ln(1 - \lambda_0).
\]

It follows that for any (\( N \)-independent) parameter variation

\[
\beta(\delta F_N - \delta \Omega^{(p)}) = \left( \frac{1}{1 - \lambda_0} - \frac{N - 1}{\lambda_0} \right) \delta \lambda.
\]
which vanishes if \( \zeta_0 \) in (49) is chosen so that
\[
N = 1/(1 - \lambda_0). \tag{60}
\]
We conclude that for this choice of \( \lambda_0 \), expectations at fixed \( N \) and in the monomer number ensemble are in fact identical to leading order. This is not a surprise. Quite generally, for fluids, (see e.g. [20]) inversion of the transformation from an \( N \)-ensemble to \( \mu \)-ensemble follows the direct prescription above to leading order, with a correction term for expectations going as \( (\delta N \gamma)^2/N^2 \). For fluids, this indeed goes as \( 1/N \), but for a chain, using (19), such as \( N_0 \), is therefore not a correction that vanishes as \( N \rightarrow \infty \). Another means of assessment is mandatory. In (12), we see that the relative correction to \( N \) has an amplitude \( \sim (1 - \lambda_0)/(1 - \lambda_1) \); the same is true for other expectations. Now suppose that, at some value of the system chemical potential \( \mu \), one has the pair \( \lambda_0, \lambda_1 \). Then if \( \mu \) is increased by \( \delta \mu \),
\[
\zeta(1/2)w(1/2) \rightarrow \exp(\beta \delta \mu)\zeta(1/2)w(1/2),
\]
so that \( \lambda_0 \rightarrow \exp(\beta \delta \mu)\lambda_0 \), \( \lambda_1 \rightarrow \exp(\beta \delta \mu)\lambda_1 \). Then, as \( \exp(\beta \delta \mu)\lambda_0 \) is made arbitrarily close to 1, \( \exp(\beta \delta \mu)\lambda_1 \) will never go beyond \( \lambda_1/\lambda_0 \), and the ratio \((1 - \lambda_0)/(1 - \lambda_1)\) indeed goes as \( 1/N \).

**Appendix 2. Fluctuations to gaussian order**

We now take a first step towards including the effect of field fluctuations on the molecular density pattern. In this initial study, we have made one major approximation and several simplifying restrictions. The approximation is of course that of mean field, or selective neglect of fluctuations. On the assumption that fluctuations are Gaussian to leading order (examples in which this is not the case are far from rare, see e.g. [23]), a correction sequence is in principle routine: we expand \( \ln Z_0 \) in (22) and (27) around \( \bar{v} \) of (29). Using \( \delta^2 \Omega_0/\delta v(\bar{r})\delta v(\bar{r}') = \delta n_0(\bar{r})/\delta v(\bar{r}') \),
\[
\Xi_0 \left[ \mu - \frac{1}{2} \phi_D - \bar{v} \right] \exp \left\{ -\frac{1}{2} \bar{v} \cdot \phi^{-1} \bar{v} \right\} = \exp \left\{ -\beta \left( \Omega_0 \left[ \mu - \frac{1}{2} \phi_D - \bar{v} \right] + \frac{1}{2} \bar{v} \cdot \phi^{-1} v - \frac{1}{2} (v - \bar{v}) \cdot \delta n_0/\delta v + \phi^{-1} (v - \bar{v}) \right) \right\}. \tag{61}
\]
It readily follows (see e.g. [24]) that the density profile is given to the next order by simply averaging the density over the Gaussian field fluctuation:
\[
n(r) = \frac{\int n_0 \mu - \frac{1}{2} \phi_D - \bar{v} - A \exp \left\{ -\frac{1}{2} (M(\bar{r}, r)/M(\bar{r}, r)) \right\} dA}{\int \exp \left\{ -\frac{1}{2} (M(\bar{r}, r)/M(\bar{r}, r)) \right\} dA}, \tag{62}
\]
where \( M(\bar{r}, r) = \phi^{-1}(r, \bar{r}) - n_0 \mu - \frac{1}{2} \phi_D - \bar{v} - A \); here, \( A \) is the field amplitude fluctuation at \( r \).

**Appendix 3. Limit of tight confinement**

We have seen, in Section 2.2, that in the absence of non-neighbour interactions, the large \( N \) monomer distribution is determined by the ‘resonance state’ \( \psi_{\lambda_0} \) satisfying
\[
(\chi^{1/2} w \chi^{1/2}) \psi_{\lambda_0} = \lambda_0 \psi_{\lambda_0}, \tag{63}
\]
where \( \lambda_{\text{max}} = \lambda_0 \sim 1 \). Explicitly, since
\[
\langle r \rangle(\chi^{-1} - w)^{-1/2}|1\rangle = \langle r \rangle(\chi^{1/2} (I - \chi^{1/2} w \chi^{1/2})^{-1} \chi^{1/2})|1\rangle \sim \frac{1}{1 - \lambda_0} \langle r \rangle(\chi^{1/2} \psi_{\lambda_0}) (\psi_{\lambda_0} \chi^{1/2})|1\rangle. \tag{64}
\]
we have under these circumstances, from (9)
\[
n(r) \sim \frac{1}{1 - \lambda_0} \psi_{\lambda_0}(r)^2, \tag{65}
\]
or since \( \psi_{\lambda_0}(r) \sim \psi_1(r) \),
\[
n(r) \sim \psi(r)^2 \frac{1}{1 - \lambda_0} = N\psi_1(r)^2. \tag{66}
\]
In other words, increasing the chain length increases the mean density uniformly, as if the floppy chain is simply winding around more under the same confinement. This uninteresting behaviour is refined by two interactions that have been ignored in getting (66). First is the stiffness of successive pair orientations, equivalent to the monomers having coupled orientational degrees of freedom, a topic that has been addressed to some extent in this format in the past and will be attended to more forcefully in the future. Second is the effect of non-next neighbour interactions, which we have here studied in a preliminary fashion. The mean field \( r(r) \) that has been encountered will also have the effect of correlating pair orientations, and will of course alter the nature of the long chain resonant state. Let us see how this works. To keep the extrapolation from (63)–(66) transparent, let us rewrite (34) (using \( \zeta = \exp(\beta \mu') \)) as,
\[
\Omega_1(\zeta, \phi) = \text{Min}_{\psi_0} \left\{ \frac{1}{2} \left( \psi - \phi \right) - 2 \ln[\phi \exp(-\phi)]^{1/2} \right\} \psi \exp(\beta \mu') \tag{67}
\]
Hence, according to (38), \( \tilde{\psi} = [\zeta^{-1} \exp(\beta \mu')]^{1/2} \psi \) is normalised, \( \langle \tilde{\psi} \rangle = 1 \) and from (39),
\[
n = N\tilde{\psi}^2. \tag{68}
\]
Near resonance is now signalled by the approximate validity of
\[
(I - [\zeta \exp(-\phi)]^{1/2} \psi \exp(-\phi)]^{1/2} \tilde{\psi} = \tilde{\lambda} \tilde{\psi}. \tag{69}
\]
with small \( \tilde{\lambda} > 0 \). Equation (50) then tells us at once that using the exact
\[
\phi^{-1} v = -n, \tag{70}
\]
we have
\[
N = 1/\tilde{\lambda}. \tag{71}
\]
There are two ways of making use of the approximation (69). Most directly, we substitute (68)–(71) into (67), obtaining
\[
\beta \Omega_1 \sim \text{Min}_{\psi_0} \left\{ \frac{1}{2} n \cdot \phi n - 2 \ln[\phi \exp(\mu)]^{1/2} [n^{1/2}] \right\}. \tag{72}
\]
This is obviously too sweeping an approximation: by using what is effectively a first order correction in the argument of a variational principle, the role of the next-neighbour \( w \) in \( \beta \Omega_1 \) has vanished. But we can pick up the next order by working instead at the ‘profile equation’ level. It is only necessary to substitute (68), (70), (71) directly into (69) to rewrite the latter as

\[
[\xi \exp(\beta \nu)]^{1/2} w [\xi \exp(-\beta \nu)]^{1/2} n^{1/2} = \left(1 - \frac{1}{N}\right) n^{1/2},
\]  

(73)

and hence as

\[
\exp\left[-\frac{\beta}{2} (u - \psi n)\right] w \exp\left[\frac{\beta}{2} (u - \psi n)\right] n^{1/2} = \left(1 - \frac{1}{N}\right) \exp(-\beta \mu'_n) n^{1/2} = \Lambda n^{1/2}
\]  

(74)

(\( \mu'_n \) is the global chemical potential). The operator in (74) has a positive kernel, hence by Jentzsch’s extension of Perron–Frobenius [25], the positive eigenfunction is unique (up to a multiplicative constant) and \( \Lambda \) is real and maximal. This leads to what is in principle a simple numerical iteration: start e.g. with \( \varphi = 0 \), compute the eigenfunction \( n^{1/2} \) and normalise it to get \( \int n(r) \, dr = N \). At the 4th stage of the \( M \)-fold iteration, replace \( \varphi \) by \( (k/M) \varphi \) and repeat the process using the current function \( n \). An alternative strategy is to parameterise \( n \) and determine the parameters by Galerkin, i.e. integrate (74) with the weight function and solve the resulting algebraic equation.

Appendix 4. Details on numerics

From preliminary simulation studies we had noticed how in the absence of a repulsive core interaction between monomers we could not converge to a stationary density. Instead the density, and therefore the total particle number, either kept on growing or shrank to zero, depending on the value of the external chemical potential. This numerical instability occurs as a result of the closeness to the resonance. In these simulations we noticed, on the other hand, that despite the fact that the density itself kept changing, the global shape of the density profile reached a steady state. We therefore decided to study the normalised density \( p(r) = n(r)/N \). Once we had formulated the problem consistently in terms of \( p \) all the numerical instabilities disappeared and we were able to determine the corresponding steady state densities and total particle numbers. The minimisation is done using a Metropolis Monte Carlo algorithm at fixed temperature, starting from a suitable ansatz for the functional form of \( p(r) \). This is one of many possible techniques, and is not optimised in any well-defined sense. We can freely impose the form of the next-neighbour potential, and therefore \( w(r, r') \), the external confining potential \( u(r) \), and the long-range interaction \( \varphi(r, r') \). The main numerical difficulty arises from the need of evaluating a functional involving six-dimensional integrals nesting three-dimensional integrals, such as

\[
\int [p(r)]^{1/2} \exp\left[\frac{\beta}{2} (\mu'(r) + N \int \varphi(r, r') p(r') \, dr')\right] w(r, r') \times \exp\left[\frac{\beta}{2} (\mu'(r') + N \int \varphi(r', r'') p(r'') \, dr'')\right] [p(r')]^{1/2} \, dr \, dr'.
\]  

(75)

It turns out it is prohibitive to try to evaluate these terms directly by the use of a simple grid. Already when using 10 discretisation points simulations are too slow, while the precision is very poor. To achieve higher accuracy in the integration, we use a Legendre–Gauss quadrature method with either six or eight points in each linear direction.

As a test system we have chosen a square well external potential with \( \mu = -\mu_0 \) inside; a harmonic n.n. potential, leading to \( w(r, r') = \exp(-C_b (|r - r'| - \rho_1)^2) \); a long-range step potential \( \varphi = C_b \theta(x - \rho_2) \), where \( \mu_0 \), \( C_b \), \( \rho_1 \) and \( \rho_2 \) are energy constants, and \( \rho_1 \) and \( \rho_2 \) are characteristic lengths. In the following we take the system size to be \( L = 1 \), \( C_b = 5 \), and \( \rho_1 = 0.2 \). As ansatz for the probability density we take the composition of three Gaussian in the three spatial directions, for a total of six parameters: \( B_x, B_y, B_z, X_0, Y_0 \) and \( Z_0 \).

\[
p(x, y, z) = A \exp[-B_x(x - X_0)^2] \exp[-B_y(y - Y_0)^2] \times \exp[-B_z(z - Z_0)^2].
\]  

(76)

At each iteration of the algorithm the normalisation of \( n \) is verified so that \( \int p(x, y, z) \, dx \, dy \, dz = 1 \), and the value of \( A \) is changed accordingly.