Semiflexible Polymers in a Random Environment

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We present using simple scaling arguments and one step replica symmetry breaking a theory for the localization of semiflexible polymers in a quenched random environment. In contrast to completely flexible polymers, localization of semiflexible polymers depends not only on the details of the disorder but also on the ease with which polymers can bend. The interplay of these two effects can lead to the delocalization of a localized polymer with an increase in either the disorder density or the stiffness. Our theory provides a general criterion for the delocalization of polymers with varying degrees of flexibility and allows us to propose a phase diagram for the highly folded (localized) states of semiflexible polymers as a function of the disorder strength and chain rigidity.

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I. INTRODUCTION

In problems concerning flexible polymers in disordered media, disorder seems to play a more important role than the conformational properties of the polymers [1-12]. The effect is usually manifested in the size of the polymer, which in a simple but rather unrealistic case of Gaussian chains in a random potential, is found to be independent of the chain length L. The radius of gyration in three dimensions simply scales as \( R \sim b / (\Delta / b^3) \), where b is a typical cut off length and plays the (undetermined) role of the segment size, and \( \Delta \) is the disorder strength at a given system volume. The problem, however, is academic mainly on two accounts: firstly, the effects due to excluded volume interactions are completely ignored, and secondly the effects of finite extensibility are not captured by the infinitely flexible Gaussian chains. It is conceivable that the inclusion of these effects, though nontrivial, can modify our present understanding of the problem.

An effort in this direction has recently been made by accounting for the effects of excluded volume interactions [12]. The results show that a chain undergoes a delocalization transition as the strength of the self-avoiding interaction is increased. Since localization is usually considered as filling up of the entropic traps — created by randomness — with polymeric materials, it is understandable that a self-avoiding chain can fill these holes up to a density close to \( \rho b^3 \sim O(1) \) (\( \rho \) is the monomer density) before delocalizing and moving to the next hole. For Gaussian chains, on the contrary, delocalization does not occur simply because they have an infinite stored length and no limit up to which the holes can be filled.

In this paper we address the second of the above mentioned problems, i.e., semiflexible polymers in a random environment. Since semiflexible polymers are finitely extensible, the theory rectifies the infinitely flexible Gaussian chain model and serves as a useful description for localization of chains with varying degrees of flexibility. In contrast to Gaussian chains, localization of semiflexible polymers depends rather strongly on the conformational properties of the polymer, i.e., the ease with which a chain can bend and eventually adopt highly packed folded states. While a polymer localizes at a certain fixed strength of the disorder, it can get delocalized with an increase in the stiffness which can be brought about by decrease in the temperature. A chain of a given fixed stiffness, on the other hand, can get delocalized with the increase in the disorder strength. The interplay of the disorder strength and bending energy is instrumental in bringing about localization of semiflexible polymers.

The following section reviews the salient features of the wormlike chain (WLC) model; it provides a general criterion for localization of a semiflexible chain in random environment using simple scaling arguments. Section III summarizes the analytical difficulties involved in treating the WLC model and discusses the Weiner model for stiff chains. Section IV presents a rigorous analysis of semiflexible chains in random media using the one step replica symmetry breaking calculations. The results are summarized in Section V. The details of the one step replica symmetry calculations are included in Appendix A.

II. MODEL AND SCALING

The configurational statistics of a semiflexible polymer is described by the wormlike chain model [13 14]. In this model, a chain is considered as a continuous curve of total length L parameterized by an arc length variable s. The variation of the unit tangent vector \( \mathbf{u}(s) = \frac{\partial \mathbf{r}(s)}{\partial s} \) at different points along the curve with respect to the arc length variable determines the energy cost of bending [15-24]. To the lowest scalar order in \( \partial \mathbf{u}(s) / \partial s \), the bending energy is given by

\[
\beta \mathcal{H} = \frac{l_p}{2} \int_0^L ds \left( \frac{\partial \mathbf{u}}{\partial s} \right)^2, \tag{1}
\]

where \( \beta \) is \((k_B T)^{-1}\) and \( l_p \) is the persistence length of the polymer. It is a measure of the distance over which bond
vectors are correlated, and hence determines the effective stiffness of the chain. In general, the correlation between the unit tangent vectors at two different points along the chain backbone decays exponentially, and is given by

\[ \langle \mathbf{u}(s) \cdot \mathbf{u}(s') \rangle = \exp(-|s - s'|/l_p). \]  

(2)

The persistence length is related to the bending energy \( \epsilon \) by \( l_p = \beta \epsilon \). If such a chain interacts with \( N \) randomly distributed obstacles, the Hamiltonian is of the form

\[ \beta H = \frac{l_p}{2} \int_0^L ds \left( \frac{\partial \mathbf{u}_s(s)}{\partial s} \right)^2 + \sum_{i=1}^N \int_0^L ds W[r(s) - r_i], \]  

(3)

where \( r(s) \) is the position vector of a segment at distance \( s \) from one end of the chain; and \( r_i \) is the position vector of the \( i \)th obstacle. \( W \) is the potential that describes the interaction between the polymer and the obstacles, the form of which is yet to be specified. If the obstacle density \( \rho(r) \) is defined by

\[ \rho(r) = \sum_{i=1}^N \delta(r - r_i), \]  

(4)

the potential can simply be written as

\[ \sum_{i=1}^N W[r(s) - r_i] = \int dr W[r(s) - r] \rho(r). \]  

(5)

At the simplest level of description, the obstacle density can be assumed to have a Gaussian distribution \[2, 3\]:

\[ P[\rho(r)] = \exp \left( -\frac{1}{2} \int dr \frac{(\rho(r) - \rho_0)^2}{\rho_0} \right), \]  

(6)

where \( \rho_0 \) is the mean obstacle density given by

\[ \rho_0 = \frac{1}{V} \int dr \rho(r). \]  

(7)

To describe the effect of the disorder, one needs to perform an average over the obstacle density. Since the average is quenched, the estimation of the free energy relies on a difficult task of computing the average of the logarithm of the partition function \( Z \). The problem is usually circumvented by introducing \( n \) independent replicas of the initial system such that

\[ \ln Z = \lim_{n \to 0} \frac{1}{n} (Z^n - 1). \]  

(8)

The replicated partition function can easily be averaged over the Gaussian random potential given by Eq. (6) to yield

\[ \langle Z^n \rangle = \int_{r(0)=0}^{r(L)=R} D[r(s)] \exp[ -\beta H_n ], \]  

(9)

where the effective Hamiltonian is given by

\[ \beta H_n = \frac{l_p}{2} \sum_{\alpha=1}^n \int_0^L ds \left( \frac{\partial \mathbf{u}_s(s)}{\partial s} \right)^2 - \frac{\rho_0}{2} \sum_{\alpha,\beta=1}^n \int_0^L ds \int_0^L ds' W[r_\alpha(s) - r_\beta(s')]. \]  

(10)

where

\[ U[r_\alpha(s) - r_\beta(s')] = \int dr W[r_\alpha(s) - r] W[r_\beta(s') - r]. \]  

(11)

If the interaction between the polymer and the obstacles is assumed to be short ranged, i.e., \( W[r] \propto \delta(r) \), then the potential can be expressed in terms of the natural scales, i.e., \( U(r) \equiv g l_p \delta(r) \). In writing the last expression for \( U(r) \), we have introduced a dimensionless coupling constant \( g \), which in general contains microscopic details. The natural (short) lengthscale defined in the problem is the persistence length \( l_p \) itself. Therefore, we can express the scaling of the potential in terms of the persistence length; this makes the second term of Eq. (10) dimensionless, as should be the case \[2\]. Having substituted this form of the potential, the effective Hamiltonian is given by

\[ \beta H_n = \frac{l_p}{2} \sum_{\alpha=1}^n \int_0^L ds \left( \frac{\partial \mathbf{u}_s(s)}{\partial s} \right)^2 - \frac{g l_p}{2} \sum_{\alpha,\beta=1}^n \int_0^L ds \int_0^L ds' \delta[r_\alpha(s) - r_\beta(s')]. \]  

(12)

where \( \Delta = gl_p \rho_0 \) and has the dimensions of length. Formally the second term in the above expression can be interpreted as a random potential which acts along the contour of the chain; it is a starting point of many studies on polymers in a disordered environment (see e.g. \[13\]). In the present study, the use of an array of obstacles instead of a random potential introduces a natural length scale in terms of the mean distance between the obstacles, \( l_{obs} \approx 1/\rho_0^{1/3} \). It turns out to be useful since this length scale can directly be compared with the persistence length in various physical situations.

Let us proceed by providing a rough estimate of the different energies involved using simple scaling arguments. This can be done by dimensional analysis of the above Hamiltonian. In the case of annealed average there is no coupling between various replicas, that is \( \alpha = \beta \), and the replica trick is not necessary. The free energy in three dimensions scales as

\[ \beta \langle \mathcal{F} \rangle \approx \frac{l_p L}{R^2} - \frac{\Delta L^2}{R^2}. \]  

(13)

The first term accounts for the bending energy penalty (per \( k_B T \)) for a chain of length \( L \). It is typically defined as the ratio of the bending force constant \( \beta \epsilon \) (= \( l_p \)) to the radius of curvature \( R \). Note that the first term in
Eq. (12) corresponds to the bending energy per thermal energy of a semiflexible chain. The derivative of the tangent vector $u(s)$ is nothing but the local curvature. The second term corresponds to excluded volume type interactions between the chain segments, which by virtue of the disorder average, are effectively attractive. In the annealed case the problem is similar to a semiflexible chain with attractive interactions. In the case of quenched disorder, on the other hand, there is a coupling between various replicas and the free energy scales as

$$\beta \langle F \rangle \simeq \frac{l_p L}{R^2} - \frac{\Delta^{1/2} L}{R^{3/2}},$$

where the second term in Eq. (14) describes a typical energy barrier of the quenched random potential; it is usually estimated as the root of the variance using simple dimensional analysis [6, 13]. It is to be noted that $\Delta = \Delta [\ln V]$, where $V$ is the volume of system. The issue of finite system volume is important for the case of quenched disorder since in an infinite random medium the disorder average, are effectively attractive. In the annealed case the problem is similar to a semiflexible chain

In what follows it is assumed that the system volume is much larger than the polymer dimensions $R^3$. The minimum of quenched free energy estimate yields the following result for the size of the chain,

$$R_{\text{loc}} \simeq l_p \left( \frac{l_p}{\Delta} \right),$$

which as in the limit of totally flexible chains, does not depend on the chain length itself but is completely determined by the disorder and the persistence length. Although the size turns out to be similar to that of a Gaussian chain, its dependence on $l_p$ instead of $b$ suggests a natural condition on the dimensionless ratio $l_p/\Delta$. That this indeed is the case can be seen from the following arguments: when the chain is reasonably stiff $l_p \approx L$ and $R_{\text{loc}}$ can at most be of the order of $L$, which immediately implies that $l_p/L \leq \Delta/l_p$; in the limit of sufficiently flexible chains, it is expected that $R_{\text{loc}} \leq R_{\text{Gauss}} (\approx \sqrt{L l_p})$, which implies $(l_p/L)^{1/2} \leq \Delta/l_p$.

In addition to the upper cut-off on the size $R_{\text{loc}}$, there is a lower cut-off, which is important for a flexible chain and suggests that $R_{\text{loc}}$ cannot be less than the persistence length $l_p$. $R_{\text{loc}} \geq l_p$ therefore implies $\Delta/l_p \leq 1$. For a flexible chain, i.e., $L/l_p > 1$, the latter constraint together with the constraint $l_p/L \leq \Delta/l_p$ can be summarized as

$$\left( \frac{l_p}{L} \right)^{1/2} \leq \frac{\Delta}{l_p} \leq 1.$$

The second inequality implies that $l_p < (g \rho_0)^{-1/3}$, which suggests that as the chain become stiffer it localizes at progressively lower density of the disorder $\rho_0$. In other words, since $l_{\text{obs}} \approx \rho_0^{-1/3}$ the chain is localized as long as its persistence length $l_p$ is less than the mean separation between the obstacles $l_{\text{obs}}$. It is to be noted that the size of the chain has a simple interpretation in terms of the obstacle density. This can be seen by substituting $\Delta \approx l_p \rho_0$ into Eq. (15) to produce $R_{\text{loc}} \propto l_p (l_{\text{obs}}/l_p)^{3}$. Then the criterion for localization, as discussed above, is simply given by the comparison between the persistence length and the mean distance between the obstacles.

As discussed in the Introduction, a Gaussian chain undergoes a continuous decrease in its size with an increase in the obstacle density since the chain can be packed without limit into the cavities formed by the obstacles. In the presence of excluded volume interactions, however, a Gaussian chain can fill this cavity only up to a point before delocalizing and moving on to the next cavity; at high obstacle density the chain eventually localizes by forming pearl necklaces (as discussed in Ref. [12]). The present theory shows that a semiflexible chain without excluded volume interactions can delocalize when the obstacle density and persistence length are related by Eq. (16). This is in marked contrast to the infinitely flexible Gaussian chain which can delocalize only when aided by the excluded volume interaction. In what follows we shall study the localization-delocalization transition of a semiflexible chain in the absence of excluded volume interactions, paying special attention to the flexible limit ($L/l_p > 1$). We will show that even in the flexible limit, new physics arises because the persistence length defines a lower cut-off length scale which is absent in the Gaussian chain. We will also compare our results in this limit to earlier results for the Gaussian chain.

Eq. (16) provides a general criterion for the delocalization of a polymer with varying degree of flexibility. The next section addresses some of the difficulties involved in treating the wormlike chain model and presents a simplified model to treat semiflexibility.

### III. Gaussian Stiff Chains

The wormlike chain model, though a proper description of semiflexible polymers, suffers from its analytical intractability in a large number of applications due to the constraint of finite extensibility, i.e., $|u(s)|^2 = 1, \forall s \in [0, L]$. In many of the practical problems, therefore, it is customary to relax the local constraint of unit tangent vector with the global constraint $\langle |u(s)|^2 \rangle = 1$ [17-24], producing a rather simple model in which the chain Hamiltonian has a connectivity term along with the bending energy term:

$$\beta \mathcal{H}_0 = \frac{\eta}{2} \int_0^L ds |u(s)|^2 + \frac{\epsilon}{2} \int_0^L ds \left( \frac{\partial u(s)}{\partial s} \right)^2.$$  

The mean square end-to-end distance obtained using the above Hamiltonian is identical with that of the wormlike chain model when $\eta = 3/2L_{p0}$ and $\epsilon = 3l_{p0}/2$ [17-24]. The correlation between the unit tangent vectors, when
calculated from the above Hamiltonian, is of the form
\[ (u(s) \cdot u(s')) = \exp(-|s - s'|/l_0), \] (18)
where \( l_0 = 2l_p/3 \). The comparison of the above Equation with Eq. (2) suggests that the persistence length for this approximate model for stiff chains is smaller by a factor of 2/3 [20, 21, 24]. This is not a serious concern in the context of the present problem since the Gaussian model contains the essence of semiflexibility, i.e., finite extensibility, while remaining analytically tractable. Moreover, the difference between \( l_p \) and \( l_0 \) becomes less important in higher dimensions [21].

As before, we begin by writing down the free energy for this simplified model using dimensional analysis or Flory-Imry-Ma arguments which includes the Wiener stretching term:
\[ \beta F \simeq \frac{R^2}{l_p^2} + \frac{l_p L}{R^2} - \left( \frac{\Delta}{R^3} \right)^{1/2} L. \] (19)

As expected, when the first term is ignored, and the free energy is minimized with respect to \( R \), one obtains \( R_{loc} = l_p^2 / \Delta \). However, the Wiener-term allows for more entropy, and it is natural to ask for perturbation around \( R_{loc} \). We look for solutions of the form
\[ R = R_{loc}(1 + \delta), \] (20)
where \( \delta \) is a small correction. When Eq. (16) is minimized with respect to \( R \), and Eq. (20) is substituted in the resulting expression, \( \delta \) turns out to be of the form
\[ \delta \propto \frac{1}{1 + (L/l_p)^2(\Delta/l_p)^4}. \] (21)

In the flexible limit of \( l_p < L \), Eq. (16) suggests \( \Delta / l_p < 1 \) and Eq. (20) reduces to
\[ R \approx R_{loc} \left( 1 - \left( \frac{\Delta}{l_p} \right)^4 \right). \] (22)

Although these scaling arguments appear to be reasonable and in accord with the physical intuition, the appropriate conditions and justifications remain to be recovered from a more rigorous analysis. The next section presents a systematic approach to this problem using Feynman variational method in replica space. Since for delocalized phases the replica symmetry breaking is no longer essential, the idea is to see if criteria for localization and delocalization can be derived from standard replica techniques using one step replica symmetry breaking.

### IV. VARIATIONAL METHOD IN REPLICA SPACE

A semiflexible polymer in presence of the randomly distributed obstacles confined in a harmonic potential is of the form
\[ \beta H = \frac{\eta}{2} \int_0^L ds \left( \frac{\partial^2 r(s)}{\partial s^2} \right)^2 + \frac{\epsilon}{2} \int_0^L ds \left( \frac{\partial r(s)}{\partial s} \right)^2 + \sum_{i=1}^N \int_0^L ds W[r(s) - r_i] + \frac{\mu}{2} \int_0^L ds r(s)^2, \] (23)
where \( \mu \) is the strength of the harmonic potential and defines the system size; the importance of this term will be discussed shortly.

Following the same steps as used to obtain Eq. (12), the quench average over the obstacle density produces the following effective Hamiltonian:
\[ \beta H_n = \frac{\eta}{2} \int_0^L ds \left( \frac{\partial r_a(s)}{\partial s} \right)^2 + \frac{\epsilon}{2} \int_0^L ds \left( \frac{\partial^2 r_a(s)}{\partial s^2} \right)^2 + \frac{\mu}{2} \int_0^L ds r_a(s)^2 - \frac{\Delta}{2} \sum_{a,b=1}^n \int_0^L ds \int_0^L ds' \delta \left[ r_a(s) - r_b(s') \right]. \] (24)

It is to be noted that carrying out an average over the disorder density produces an effective attractive interaction of strength \( \Delta \) between different replicas. To estimate the average free energy we employ the Feynman variational method in replica space [21–23], which is of the form
\[ n \langle F \rangle = \langle H_n - h_n \rangle_h_n - \frac{1}{\beta} \int \Pi_d a \mathcal{D}[r_a(s)] e^{-\beta h_n}, \] (25)
where \( h_n \) is the Gaussian trial Hamiltonian with variational function \( \sigma(s - s') \) characterizing the strength of the harmonic potential:
\[ \beta h_n = \frac{\eta}{2} \sum_{a=1}^n \int_0^L ds \left( \frac{\partial r_a(s)}{\partial s} \right)^2 + \frac{\epsilon}{2} \sum_{a=1}^n \int_0^L ds \left( \frac{\partial^2 r_a(s)}{\partial s^2} \right)^2 + \mu \sum_{a=1}^n \int_0^L ds r_a(s)^2 - \sum_{a,b=1}^n \int_0^L ds \int_0^L ds' \sigma_{ab}(s - s') r_a(s) \cdot r_b(s). \] (26)

Defining the matrix of correlations by
\[ \langle r_a(s) \cdot r_b(s') \rangle = \frac{3}{L} \sum_q e^{-iq(s-s')} G_{ab}(q), \] (27)
and substituting into Eqs. (21), (25) and (26), the free energy simplifies to
\[ \frac{\beta \langle F \rangle}{L} = \text{const} + \frac{3}{2nL} \sum_{a=1}^n \sum_q (eq^4 + \eta q^2 + \mu) G_{aa}(q) - \frac{3}{2nL} \sum_q Tr \ln[G(q)] \]
where delta function of Eq. (24) has been exponentiated and averaged over the Gaussian random potential to produce the last two terms in the above equation. \( G_{ab}(q) \) is the propagator with respect to the trial Hamiltonian \( h(R) \):

\[
G_{ab}(q) = [(\epsilon q^4 + \eta q^2 + \mu)\delta_{ab} - \sigma_{ab}(q)]^{-1}.
\]

(29)

\( \sigma_{ab}(q) \), in general, is an \( n \times n \) matrix, which in the case of replica symmetric solution can be represented by a simple variational ansatz:

\[
\sigma_{ab}(q) = -\delta_{ab}[\sigma_d^2 \delta_{q,0} + \sigma_d(1 - \delta_{q,0})] + \sigma(1 - \delta_{ab})\delta_{q,0},
\]

(30)

where the diagonal variational parameters \( \sigma_d \)'s representing the \( q = 0 \) modes have been separated from all other modes (represented by \( \sigma(q) \)) to avoid working with infinite number of variational parameters. The off-diagonal terms have been considered to depend only on the \( q = 0 \) mode, as is conventionally done in \textsuperscript{11}. Translational invariance of the variational function requires \( \sum_q \sigma_{ab}(q = 0) = 0 \), resulting in a condition \( \sigma_d + \sigma = 0 \) that leaves only two of the variational parameters independent.

It has been argued by Cates and Ball that in the thermodynamic limit of infinite system size the annealed and quenched average should be the same, and an infinite Gaussian chain should always be collapsed \textsuperscript{4}. That this indeed is the case has been shown by Goldshmidt, where the \( \mu \to 0 \) limit corresponds to an infinite system size, and the replica symmetric solution for a quenched random potential recovers the results of the annealed disorder. On the contrary, when \( \mu \) is taken to be finite, the replica symmetric solution is found to be an inadequate description of the problem, especially at high disorder strength where replica symmetry needs to be broken \textsuperscript{11}. The importance of weak replica symmetry breaking was recognized by one of us earlier \textsuperscript{10}, where a special field theoretic description of polymers in strong disorder was used. In general, for problems concerning random potentials with short range correlations, one step replica symmetry breaking has been found to be a sufficient description \textsuperscript{25}. In what follows, therefore, we employ one step replica symmetry breaking to estimate the size of a semiflexible polymer in a quenched random distribution of obstacles.

From Eq. (27), the mean square end-to-end distance of a polymer in terms of the propagator \( G(q) \) is given by

\[
\langle R^2 \rangle = \lim_{n \to 0} \frac{3}{n} \sum_{a,b=1}^n \int_0^L \frac{dk}{2\pi} e^{-k^2} \sum_q (G_{aa}(q)+G_{bb}(q)-2G_{ab}(q)e^{-iqL}),
\]

(31)

where in the limit of large \( L \) the summation over \( q \) has been replaced by an integral. Eqs. (29) and (30) when substituted into the above equation followed by integration over \( q \) produces

\[
\langle R^2 \rangle = \frac{3}{(\eta + 2\sigma_d^2/\epsilon^{1/2})} \frac{1}{\sigma_d^{1/2}} \left( \frac{1}{\alpha_+^{1/2}} - \frac{1}{\alpha_-^{1/2}} \right)
\]

(32)

where \( \alpha_{\pm} = [1 \pm (4\epsilon \sigma_d/\eta)^{1/2}]^{1/2} \) and \( \sigma_d \) is the variational parameter to be determined by minimizing the variational free energy within one step replica breaking scheme \textsuperscript{11,25}.

In replica symmetry breaking scheme, the \( n \times n \) matrix \( G(q) \) in the limit of \( n \to 0 \) is parameterized by a function \( g(q,u) \), where \( u \) is a continuous variable that is defined in the interval \([0,1]\). The diagonal elements, on the other hand, are parameterized by \( \tilde{g}(q) \) such that

\[
\lim_{n \to 0} \frac{1}{n} \sum_{a,b=1}^n G_{ab}(q) = \tilde{g}(q) - \langle g(q) \rangle,
\]

(33)

where \( \langle g(q) \rangle = \int_0^1 du g(q,u) \).

The idea is to compute the inverse of a hierarchical matrix \( \tilde{H}(q) = [(\epsilon q^4 + \eta q^2 + \mu)\tilde{\delta}_{ab} - \tilde{\sigma}_{ab}(q)] \) such that \( G(q) = \tilde{H}(q)^{-1} \). The matrix \( \tilde{H}(q) \), in the limit of \( n \to 0 \), is similarly parameterized by the diagonal element \( \tilde{h}(q) \) and off-diagonal element \( h(q,u) \) given by

\[
h(q,u) = \epsilon q^4 + \eta q^2 + \sigma_d \delta_{q,0} + (1 - \delta_{q,0})\sigma_d - \sigma(q,u),
\]

(34)

In one step replica symmetry breaking, the symmetry is broken at a point \( u_c \), and the function is defined by

\[
\sigma(q,u) = \begin{cases} 
\sigma_0 & u < u_c, \\
\sigma_1 & u > u_c.
\end{cases}
\]

(35)

Translational invariance of the variational function in the limit of \( n \to 0 \) for the case of one step replica symmetry breaking requires \( \sigma_d + u_c\sigma_0 + (1 - u_c)\sigma_1 = 0 \). The variational free energy, as given by Eq. (28), in the limit of \( n \to 0 \) is therefore of the form:

\[
\frac{\beta \langle F \rangle}{L} = \text{const} + \frac{3}{2} \int \frac{dq}{2\pi} (\epsilon q^4 + \eta q^2 + \mu)\tilde{g}(q) + \lim_{n \to 0} \frac{3}{2n} \int \frac{dq}{2\pi} \text{Tr} \ln[H(q)]
\]

\[
-\frac{\Delta}{2} \int_0^L dx \int \frac{dk}{2\pi} e^{-k^2} \int \frac{dq}{(2\pi)^3} e^{-i(qL)} \tilde{g}(q)(1 - e^{-iqx})
\]

\[
-\frac{\Delta}{2} \int_0^1 du \int_0^L dx \int \frac{dk}{(2\pi)^3} e^{-k^2} \int \frac{dq}{(2\pi)^3} e^{-i(qL-2\pi)} \tilde{g}(q) - g(q,u) e^{-iqx}.
\]

(36)

The details of the replica symmetry breaking (RSB) method are described in the paper by Mézard and Parisi.
of ∆

the appendix of which provides the explicit expressions for some of the terms introduced in the above equation. One step replica symmetry breaking steps relevant to the present problem are furnished in Appendix A. The free energy as given by Eq. (A14), in the limit of large L and μ → 0, is dominated by

\[ \frac{\beta \langle F \rangle}{L} = \text{const} - \frac{3a_d g_0}{4} + \frac{3}{2g_0 \epsilon^{1/2} \sigma_d^{1/2}} - \tilde{\Delta}^{1/2} g_0^{-3/4}, \]

(37)

where \( g_0 = (\eta + 2\sigma_d^{1/2} \epsilon^{1/2} - 1/2) \frac{1}{\sigma_d^{1/2} \epsilon^{1/2}} \) and \( \tilde{\Delta} = 3\Delta |\ln \mu|/(2\pi)^{3/2} \). The minimization of the free energy with respect to \( \sigma_d \) produces the following algebraic equation:

\[ \tilde{\Delta}^4 (\eta + 2\sigma_d^{1/2} \epsilon^{1/2})^2 - \sigma_d = 0. \]

(38)

That Eq. (39) is a valid description in both stiff and flexible limit can be seen as follows: In the absence of disorder (corresponding to \( \sigma_d \to 0 \)), when \( L \eta l^{1/2}/\epsilon^{1/2} \gg 1 \), \( \langle R^2 \rangle \approx 3L \eta (\approx 2L \eta p) \) (the flexible limit); when \( \eta^{1/2} L/\epsilon^{1/2} \ll 1 \), \( \langle R^2 \rangle \approx L^2 \) (the rod limit).

V. RESULTS AND DISCUSSION

A. Localization-Delocalization Transition

When \( \eta = 3/2l_p \) and \( \epsilon = 3l_p/2 \) are substituted into Eq. (38), it can be rewritten in the following dimensionless form:

\[ \Delta_0^4 (1 + 3\sigma_0^{1/2})^7 - \sigma_0 = 0, \]

(39)

where \( \Delta_0 = \tilde{\Delta}/l_0 \), \( \sigma_0 = \sigma_d l_0^3 \), and \( l_0 = 2l_p/3 \).

The range of applicability of one-step RSB solution can qualitatively be seen from the behavior of \( u_c \), which is defined in the interval [0, 1]. The form of \( u_c \) as given by Eq. (A11), suggests that \( |\ln \mu| \leq L \Delta_0^{1/2}/g_0^{3/4} \), where \( g_0 = l_0^2 (1 + 3\sigma_0)^{-1/2} \sigma_0^{-1/2} \).

The above inequality implies that a chain localizes when the binding energy is greater than the translational energy. It is to be noted that this inequality arises as a natural condition on the validity of replica symmetry breaking solution. In the flexible limit of \( L/l_0 > 1 \) and \( l_0 \to 0 \), Eq. (39) suggests that \( \sigma_0 \approx (\Delta_0/l_0)^4 \), and the above inequality yields \( (l_0/L)^{1/2} |\ln \mu|^{1/2} \leq \Delta_0/l_0 \), the form of which is same as suggested using simple scaling arguments after Eq. (15).

Eq. (39) can be solved numerically for \( \sigma_0 \) as a function of \( \Delta_0 \). The results are summarized in Fig. (1); the localized region corresponds to the case where one-step RSB does not provide physically meaningful solutions to the dimensionless variational parameter \( \sigma_0 \). As is evident from Fig. (1), the chain is localized as long as the condition \( \Delta_0/l_0 \leq 1 \) is satisfied. The latter condition introduces an upper cut-off on the dimensionless ratio \( \Delta_0/l_0 \); it suggests that in the flexible limit \((0 < l_p/L < 1)\), for a given persistence length, there exist a critical disorder strength \( \Delta_c(< l_0) \) above which the chain delocalizes.

The Gaussian approximation for semiflexible chains forbids us to consider rigorously the localization of a rigid rod [17-21]. In what follows, we restrict ourselves to the limit of intermediate stiffness \( 0 < l_p/L < 1 \). The real stiff limit of \( L/l_0 < 1 \) requires a separate treatment.

B. The size of a semiflexible polymer in disorder

In the flexible limit of \( l_0 \to 0 \), Eq. (39) can be solved to quadratic order in \( \sigma_0 \). The resulting expression is given by

\[ \sigma_0^{1/2} = \frac{\Delta_0^4 [1 + (12(1 - \Delta_0^4)/7\Delta_0^{4/1})^{1/2}]/2}{(1 - \Delta_0^4)}. \]

(40)

When \( \eta = 3/2l_p \) and \( \epsilon = 3l_p/2 \) are substituted into Eq. (31), the resulting expression can be written in the following form:

\[ \langle R^2 \rangle = \frac{3l_p^2}{(1 + 3\sigma_0^{1/2})^{1/2} \sigma_0^{1/2}} \left( 1 - \frac{1}{\alpha_{1/2} - \alpha_{-1/2}} \right) \times \left( \alpha_{1/2} e^{-2\alpha_{1/2} L/3l_0} - \alpha_{-1/2} e^{-2\alpha_{1/2} L/3l_0} \right) \]

(41)

where \( \alpha_{\pm} = [1 \pm (1 - 9\sigma_0^{1/2})/2] \). In the flexible limit of \( L/l_0 > 1 \), to leading order in \( l_0 \), the above expression can be simplified to produce the following approximate expression for the mean square end-to-end distance:

\[ \langle R^2 \rangle = \frac{3l_p^2}{\sigma_0^{1/2}} (1 - \exp(-\sigma_0^{1/2} L/l_0)). \]

(42)
In the limit of low disorder strength $\Delta_0 \to 0$, $R \approx l_p^3/\bar{\Delta}$, which is same as Eq. (11) obtained using scaling arguments. When the disorder strength reaches its critical value corresponding to $\Delta_0 \approx 1$, $R \approx l_p^3/\bar{\Delta}^2$. The latter suggests that when the average separation between the obstacles is of the order of the persistence length ($l_{obs} \approx \rho_0^{-1/3} \approx l_0$), the disorder can explore the length scale over which bending energy is stored. Since any further increase in the disorder density amounts to high energy penalty for bending, the polymer delocalizes above the critical density. The decrease in the size close to $\Delta/l_0 \approx 1$ can therefore be attributed to the dominance of the bending mode at this length scale. The results are summarized quantitatively in Fig. (2), which is a numerical solution of Eqs. (39) and (41) for the ratio $L/l_0 = 100.0$.

It is to be noted that in the above analysis we have used $l_0$ instead of $l_p$ as the persistence length. This is because $l_0 = 2l_p/3$ appears as the natural persistence length for Gaussian stiff chains, an issue that has been discussed in Sec. III. In the rest of our analysis $l_p$ will be used as the persistence length.

C. The final state of the polymer

When the polymer is able to find a suitable region among the obstacles, its conformation must contain a rich structure. Though the results of the previous section show a dramatic decrease in the chain size with the increase in the disorder strength, they say nothing about several different folded states a chain can adopt as a function of the chain rigidity $L/l_p$ and dimensionless disorder strength $\bar{\Delta}/l_p$. In what follows we propose a phase diagram for several possible compact states of the localized polymer by comparing their surface energies.

Since an average over the disorder density produces an effective attractive interaction between chain segments, a chain can find itself in several compact states depending on its stiffness. For a fully flexible chain the compact state is expected to be a globule; a stiffer chain, on the other hand, has a tendency to wind around itself many times to form a toroid [24,27]. In presence of obstacles, the size of a circular toroid can be obtained by minimizing the free energy of the form

$$\beta F = \frac{l_p R^2}{R_0 b} - \left( \frac{\bar{\Delta}}{R} \right)^{1/2} \frac{l_p^3}{R^2},$$

where $b$ is the monomer size and $l_p$ is the length over which monomers of size $b$ are correlated. The first term accounts for the bending energy of a toroid $R_0$, $R_s$ is the radius of the torus tube, and $R$ is the radius from the center of the hole to the center of the tube. Minimization of the free energy with respect to $R$ yields $R \approx l_p^3/\bar{\Delta}$ and $R_s \approx \bar{\Delta}^{1/2}L^{1/2}l_p/b$, where the volume constraint, $V = 2\pi^2 R R_s^2 = \pi b^2$, has been used. As expected, the minor radius $R_s$ grows at the expense of the major radius $R$ with the decrease in the persistence length implying that for a given disorder strength there exist a critical ratio of $L_{crit}/l_p$ above which the chain prefers to be in a globular state. Comparison of the surface energy penalty of a toroid $S_t \approx \gamma R R_s$ with respect to a sphere $S_g \approx \gamma R^2$ suggests $L_{crit}/l_p \approx (l_p/\bar{\Delta})^3(l_p/b)^2$, where $\gamma$ is the surface tension. As expected, for high disorder strength, $\bar{\Delta}/l_p$, and, $L/l_p > L_{crit}/l_p$, the chain prefers to be in a globular state. In the opposite limit the chain adopts a toroidal state.

When the disorder strength is high, a chain can pack densely only if it chooses a mean direction that a given distribution of obstacles imposes. In other words, a high obstacle density favors anisotropy in structures, and it seems unlikely that under these conditions a polymer can adopt uniformly bent states like globules or toroids. It is physically intuitive, however, that if the polymer is able to localize for moderate persistence lengths, it can do so by producing many hairpins in a small available volume. The hairpins are the abrupt reversals in the direction of the chain from the mean ordering direction and result from the balance between the elastic penalty that favors gradual bends, and orientational field that favors rapid
VI. CONCLUSIONS

We interpret our results to suggest that the interplay between the two length scales — the persistence length of the chain and the average separation between the obstacles — serves to localize a semiflexible polymer in a random distribution of obstacles. As long as the disorder density is low, the mean separation between the obstacles is larger than the persistence length, and the chain is localized; but as soon as the disorder explores the length scales of the order of the persistence length the chain delocalizes since the energy penalty for bending exceeds the energy gain in binding. On the contrary, when the density of obstacles is kept fixed, delocalization can be brought about by varying the temperature. While the ease of localization of a semiflexible chain can be enhanced by increase in the temperature (corresponding to the decrease in the persistence length), a localized flexible chain can be delocalized by lowering the temperature. Once the polymer is localized it adopts a highly compact folded state by forming hairpins for moderate persistence length and high disorder strength; for small and large persistence length the chain conformation is dominated by uniformly bent shapes like globules and toroids respectively.

Although the idea was to present a general theory for localization of semiflexible polymers, there is one respect in which the present formalism could be seen to have limitations, and that is the use of the Gaussian approximation for stiff chains, which fails to adequately describe the rigid rod limit. Nevertheless, given the range of systems that can be described by flexible chain models, this theory holds out the possibility of addressing several realistic problems, for instance, the immobilization of biopolymers on surfaces with random interactions.
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APPENDIX A: ONE-STEP REPLICA SYMMETRY BREAKING

The definitions of some of the terms introduced in Eq. (36) is as follows:

\[ \lim_{n \to 0} \frac{1}{n} Tr \ln[H(q)] = \ln(\tilde{h}(q) - \langle h(q) \rangle) \]

\[ + \frac{h(q,0)}{h(q) - \langle h(q) \rangle} \int_0^1 \frac{du}{u^2} \ln \frac{\tilde{h}(q) - \langle h(q) \rangle - [h](q,u)}{h(q)} \]

\[ \tilde{g}(q) = \frac{1}{h(q) - \langle h(q) \rangle} \left[ 1 - \frac{1}{u^2} \frac{[h](q,u)}{h(q) - \langle h(q) \rangle - [h](q,u)} \right] \]

\[ g(q,u) = - \frac{1}{h(q) - \langle h(q) \rangle} \left[ \frac{u[h](q,u) - \langle h(q) \rangle - [h](q,u)}{[h](q,u)} \right] \]

\[ e \int u \frac{\tilde{h}(q) - \langle h(q) \rangle - [h](q,u)}{[h](q,u)} \]

For the case of one step replica symmetry breaking the calculation is straightforward, and the explicit expressions for the above equations are given by

\[ \hat{h}(q) - \langle h(q) \rangle = e q^4 + \eta q^2 + \mu + \sigma(1 - \delta_{\alpha,0}), \]

\[ \tilde{g}(q) = \delta_{\alpha,0} \left[ \frac{1}{\mu} + \frac{\alpha_0}{\mu^2} - \left( \frac{1}{u_\alpha} \right) \frac{\Sigma_1}{\mu(\mu + \Sigma_1)} \right] \]

\[ + \frac{(1 - \delta_{\alpha,0})}{(eq^4 + \eta q^2 + \sigma_0)} \]

\[ g(q,u < u_c) = \delta_{\alpha,0} \frac{\sigma_0}{\mu^2} \]

\[ g(q,u > u_c) = \delta_{\alpha,0} \left[ \frac{\Sigma_1}{u_c \mu(\mu + \Sigma_1)} + \frac{\sigma_0}{\mu^2} \right], \]

with

\[ [h](q,u) = 0; \quad u < u_c, \]

\[ = -\Sigma_1 \delta_{\alpha,0}; \quad u > u_c, \]

where \( \Sigma_1 = u_c(\sigma_1 - \sigma_0) \). Having substituted the above equations into Eq. (36) followed by integrations with respect to the variables \( q \) and \( k \), the expression for the free energy simplifies to

\[ \frac{\beta \langle \mathcal{F} \rangle}{L} = const + \frac{3(\mu - \sigma_d)g_0}{4} + \frac{3}{2g_0(\sigma_d)^{1/2}} \]

\[ + \frac{3}{2L} \left[ \ln \frac{\mu}{\sigma_d} - \frac{\mu}{\sigma_d} \right] \left( 1 - \frac{1}{u_c} \right) \]

\[ \times \left( \frac{\mu + \Sigma_1}{\mu} \frac{\Sigma_1}{\mu + \Sigma_1} \right) \]

\[ - \frac{\Delta L}{2(2\pi)^{3/2} g_0^{3/2}} \left( \int_0^1 dx (g_1(x))^{-3/2} \right. \]

\[ \left. - g(u > u_c)^{-3/2} - g_c(g(u < u_c)^{-3/2}) \right), \]

where \( g_0 = (\eta + 2\sigma_d^{1/2}e^{1/2} - 1/2\sigma_d^{-1/2} \); other terms introduced in the above expression have the following definitions:

\[ g(u > u_c) = 1 + \frac{2}{L g_0} \left( \frac{1}{\mu + \Sigma_1} - \frac{1}{\sigma_d} \right), \]

\[ g(u < u_c) = 1 + \frac{2}{L g_0} \left( \frac{1}{\mu + \Sigma_1} - \frac{1}{\sigma_d} \right) + \frac{\Sigma_1}{u_c \mu(\mu + \Sigma_1)}, \]

\[ g_1(x) = 1 - \frac{1}{\alpha_+ e^{1/2} - \alpha_+ e^{-1/2} g_1(x)^{-3/2} - \alpha_- e^{-1/2} g_1(x)^{-3/2}}, \]

where \( \alpha_\pm = [1 \pm (1 - 4e\sigma_d/\eta^2)^{1/2})]/2 \). The free energy can be minimized with respect to \( \Sigma_1 \) and \( u_c \). In the limit of large \( L \) and \( \mu \to 0 \), the solution of the resulting equations yield

\[ \Sigma_1 = \Delta_1^{1/2} g_0^{-7/4}, \]

\[ u_c = 3g_0^{3/4} |\ln \mu| \Delta_1^{1/2} L^{-1}, \]

where \( \Delta = 3\Delta |\ln \mu|/(2\pi)^{3/2} \). The substitution of above equations into Eq. (36) implies the expression for the free energy to

\[ \frac{\beta \langle \mathcal{F} \rangle}{L} = \frac{3(\mu - \sigma_d)g_0}{4} + \frac{3}{2g_0(\sigma_d)^{1/2}} - \Delta_1^{1/2} g_0^{-3/4} \]

\[ + \frac{\Delta_0^{-3/2} L}{6 |\ln \mu|} \int_0^1 dx (g_1(x))^{-3/2} - 1 \]

\[ + \frac{\Delta_0^{-5/2}}{2 |\ln \mu|} (\Delta_1^{1/2} g_0^{7/4} - \sigma_d^{-1}), \]

where only terms of \( O(1) \) with respect to expansion in \( 1/L \) have been retained.
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