Radial distribution function of semiflexible polymers

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We calculate the distribution function of the end–to–end distance of a semiflexible polymer with large bending rigidity. This quantity is directly observable in experiments on single semiflexible polymers (e.g., DNA, actin) and relevant to their interpretation. It is also an important starting point for analyzing the behavior of more complex systems such as networks and solutions of semiflexible polymers. To estimate the validity of the obtained analytical expressions, we also determine the distribution function numerically using Monte Carlo simulation and find good quantitative agreement.

While we have a comparably complete theoretical picture of highly flexible chain molecules, the statistical mechanics of semiflexible polymers is a field with a number of open questions that has received renewed attention lately. Considerable motivation stems from the crucial importance of the elasticity of biopolymers like spectrin, actin, and microtubules for the mechanical properties of cells. Recent advances in visualizing and manipulating such macromolecules have provided unique experimental tools for the study of the static and dynamic properties of single filaments. A quantitative measure for a polymer’s flexibility is its persistence length \( \ell_p \), which is the characteristic length governing the decay of tangent–tangent correlations. Nature provides polymers of very different stiffness, e.g., \( \ell_p \approx 10 \text{ nm} \) for spectrin, \( \ell_p \approx 50 \text{ nm} \) for DNA, \( \ell_p \approx 17 \text{ \mu m} \) for actin, and \( \ell_p \approx 5.2 \text{ nm} \) for microtubules. On length scales larger than approximately \( \ell_p \) a polymer with contour length \( L \gg \ell_p \) (flexible polymer) can be described as a self–avoiding freely jointed chain. For molecules with \( t := L/\ell_p \) of the order of one (semiflexible polymer) this is not possible, as can be seen immediately by comparing a typical contour (e., from Ref. 3) to the random walk corresponding to a freely jointed chain. Even for long strands of DNA with \( t \approx 1000 \) measurements of the molecule’s extension at large forces have revealed significant deviations from the freely jointed chain model, which find their explanation in bending elasticity. Thus it is essential to consider models which take chain rigidity into account. For sufficiently stiff chains effects resulting from self–avoidance can be neglected due to the strong energetic suppression of configurations where the chain folds back onto itself. Furthermore the polymers under consideration can be regarded as inextensible. The corresponding model is the wormlike chain introduced by Kratky and Porod almost 50 years ago.

A central quantity for characterizing the conformations of single polymer chains is the distribution function \( G(\mathbf{r}; L) \) of the end–to–end distance \( \mathbf{r} \) for given contour length \( L \) and persistence length \( \ell_p \). For models like the wormlike chain with only short–range interactions between monomers, \( G(\mathbf{r}; L) \) actually is the probability density of finding any two monomers at relative position \( \mathbf{r} = \mathbf{r}(s) - \mathbf{r}(s') \) where \( L = |s - s'| \) is the distance between the monomers along the chain. For a freely jointed phantom chain \( G(\mathbf{r}; L) \) is known exactly and for many purposes is well approximated by a simple Gaussian. Complications arise when the self–avoidance of real chains is taken into account. In previous investigations of the wormlike chain \( G(\mathbf{r}; L) \) was obtained approximately for almost fully flexible polymers (large \( t \)) in the form of corrections to the Gaussian distribution function up to order \( t^{-2} \) \( \text{(13,14)} \). For general \( t \) only the lowest three even moments were calculated analytically. Higher even moments were obtained by numerical techniques. For polymers close to the stiff limit all even moments of the distribution function were calculated in an expansion in \( t \) \( \text{(3,17)} \), but the expressions obtained for \( G(\mathbf{r}; L) \) in this limit are only given up to quadratures \( \text{(17,18)} \) and do not show the correct qualitative behavior when integrated numerically.

In this letter we determine \( G(\mathbf{r}; L) \) in two– and three–dimensional embedding space in an approximation valid for small \( t \) and compare the analytical expressions obtained to data from a Monte Carlo simulation. The range of validity of our results almost extends to values of the bending rigidity where the Daniels approximation \( \text{(3,14)} \) becomes applicable.

For our analytical calculations we adopt a continuum version of the wormlike chain model where the polymer is represented by a differentiable space curve \( \mathbf{r}(s) \) of length \( L \) parameterized to arc length \( s \). Its statistical properties are determined by an effective free energy

\[
\mathcal{H} = \frac{\kappa}{2} \int_0^L ds \left[ \frac{\partial \mathbf{t}(s)}{\partial s} \right]^2,
\]

where \( \mathbf{t}(s) = \partial \mathbf{r}(s)/\partial s \) is the tangent vector at arc length \( s \). The resulting persistence lengths are \( \ell_p = \kappa/k_BT \) for \( d = 3 \) and \( \ell_p = 2\kappa/k_BT \) for \( d = 2 \), where \( d \) is the dimension of the embedding space. The inextensibility of the chain is expressed by the local constraint \( |\mathbf{t}(s)| = 1 \) which leads to non-Gaussian path integrals. Note that this local curvature model is equivalent to a one-dimensional nonlinear \( \sigma \)-model. While a few quantities like \( \langle R^2 \rangle \) and \( \langle R^4 \rangle \) can be obtained exactly, one has to resort to some approximative scheme to calculate the
end–to–end distribution function
\[ G(r; L) = \langle \delta(r - R) \rangle , \]  
(2)
where \( R := r(L) - r(0) \) and \( \delta(r) \) is the Dirac \( \delta \)-function.

In order to understand the effect of possible approximations to Eq. (2) for stiff polymers, it is instructive to recapitulate the classical problem of bending a rigid rod. The energy of a straight rod of length \( L \) and bending modulus \( \kappa \) is an almost linear function of its end-to-end distance \( r \): \( E_{el} \approx f_{\kappa}(L - r) \) where \( f_{\kappa} = \kappa \pi^2 / L^2 \) is the critical force for the onset of the Euler instability (since the radial distribution function does not specify the direction of the tangent vectors at \( s = 0 \) and \( s = L \), the appropriate boundary conditions are open ends). Neglecting fluctuations around the classical contour this would lead to an end–to–end distribution function with maximum weight at \( r = L \), \( G(r; L) \propto \exp[-f_{\kappa}(L - r)/(k_BT)] \). But, for a completely stretched chain there is up to global rotations only one possible configuration and consequently the end–to–end distribution function has to vanish at full extension. Hence it is essential to take into account entropic effects. While stiff chains are energy dominated, more flexible chains are mostly governed by the entropic effects with the limit being the freely jointed chain. Approaching full extension, however, \( G(r; L) \) must vanish for all chains. This shows that while \( G(r; L) \) tends to the distribution \( \delta(L - r)/4\pi L^2 \) (\( d = 3 \)) for \( \kappa \to \infty \) and \( t \to 0 \), it can never be expanded in \( t \) around this limit (consider the probability of full extension). For this reason, it is impossible to obtain the distribution function from an expansion of all even moments in terms of \( t \). Softening the constraint of fixed contour length affects both energetic and entropic contributions to the distribution function in essential ways. If it is relaxed to the point of fixing only \( \int_0^L ds \alpha^2(s) \) by means of a single Lagrange multiplier (e.g., Ref. [21]), the distribution functions obtained are essentially Gaussian and will not show the correct qualitative behavior for stiff polymers. Failure to reproduce the vanishing of \( G(r; L) \) near full extension also results when approximations are used which neglect essential parts of the fluctuational contributions as in Ref. [22].

For end–to–end distances \( r \) close enough to full stretching and/or large values of \( \kappa \), the typical configuration of the chain will be close to a straight rod. Thus the deviations of the tangent vectors from the average direction can be treated as small variables. For \( d = 3 \) we parameterize the contour through the tangent field: \( t(s) = (a_x(s), a_y(s), 1)/\sqrt{1 + a_x^2(s) + a_y^2(s)} \), which properly takes into account the constraint of inextensibility. We employ a harmonic approximation and keep only terms up to second order in \( \mathcal{H} \), the measure factor, and the arguments of the \( \delta \)-function in Eq. (2). The error caused by this approximation vanishes near full extension. From here on we shall measure all lengths in units of \( L \) and all energies in units of \( k_BT \). With this convention we have (for \( d = 3 \)) \( \kappa = \ell_p^2 \) and \( t = \kappa^{-1} \). We also drop the second argument in \( G(r; L) \) and make rotation invariance explicit by writing \( G(r) \). Use of \( 2\pi \delta(x) = \int dq \exp(iqx) \), expansion of \( a_x(s) \) and \( a_y(s) \) in cosine series as appropriate for the boundary conditions of open ends, and evaluation of the resulting Gaussian integrals leads to
\[ G(r) = \frac{2\kappa}{4\pi N} \sum_{k=1}^{\infty} \pi^2 k^2 (-1)^{k+1} e^{-\pi^2 k^2 (1-r)} , \]  
(3)
where \( N \) is a normalization factor compensating the failure of the approximation used to conserve the normalization of \( G(r) \) for finite \( \kappa \). Details of the calculation can be found in a forthcoming publication [22]. For \( \kappa(1-r) \gtrsim 0.2 \) the distribution function is dominated by the \( \kappa = 1 \) term which is just our heuristic result from above (“Euler instability”) if the dimensional factors are put back in. For \( r \to 1 \), however, more and more terms have to be considered. By writing Eq. (3) as an integral over Fourier expanded \( \delta \)-functions it is possible to transform Eq. (3) to
\[ G(r) = \frac{1}{4\pi N} \frac{\kappa}{2\sqrt{\pi}} \sum_{\ell=1}^{\infty} \frac{1}{(\kappa(1-r))^{3/2}} \exp \left[-\frac{(\ell - 1/2)^2}{(\kappa(1-r))} \right] H_2 \left[ \frac{\ell - 1/2}{\sqrt{\kappa(1-r)}} \right] , \]  
(4)
where \( H_2(x) = 4x^2 - 2 \) is the second Hermite polynomial. This series converges very quickly for \( \kappa(1-r) \lesssim 0.2 \) where the behavior of \( G(r) \) is completely dominated by the \( \ell = 1 \) term.

In recent experiments as well as in simulations the polymer was effectively restricted to a \( d = 2 \) embedding space [1][2][23]. In this case it is convenient to choose a parameterization \( t(s) = (\cos \phi, \sin \phi) \). The resulting effective free energy is quadratic in \( \phi \), \( \mathcal{H} = (\kappa/2) \int ds (\partial \phi / \partial s)^2 \). In order to calculate \( G(r) \) we again approximate the constraints given by the \( \delta \)-functions in Eq. (2) by \( 0 = r_\ell (1) \approx \int ds \phi \) and \( r = r_\ell (1) \approx 1 - \frac{1}{2} \int ds \partial \phi^2 \). A calculation along similar lines as before [22] leads to the result
\[ G(r) = \frac{1}{2\pi N} \frac{\kappa}{\sqrt{\pi}} \sum_{\ell=0}^{\infty} \frac{(2\ell - 1)!!}{2\ell!} \frac{1}{(2\kappa(1-r))^{5/2}} \exp \left[-\frac{(\ell + 1/4)^2}{2\kappa(1-r)} \right] D_{3/2} \left[ \frac{2\ell + 1/4}{\sqrt{2\kappa(1-r)}} \right] , \]  
(5)
with \( D_{3/2}(x) \) a parabolic cylinder function. The convergence properties of Eq. (5) are similar to those of Eq. (4).

In order to assess the quality of our approximations, we have used Monte Carlo simulation to evaluate \( G(r) \) numerically. We adopted the following discretized version of the wormlike chain: The polymer is described as a chain composed of \( N \) tethers of fixed length \( a = 1/N \) and direction \( t \) with a bending energy \( \mathcal{H}_b = k a^{-1} \sum_{i=1}^{N-1} t_i \cdot t_{i+1} \) [23]. The standard Metropolis algorithm was used to measure \( G(r) \). We found that results cease to depend appreciably on \( N \) as soon as there are three to four segments in one persistence length. On the order of \( 10^6 \) MC-steps per segment were performed. Final results were obtained by averaging over several independent runs. The
The accuracy of $\langle R \rangle$ obtained was typically of the order of 0.5%. Measured expectation values $\langle R^2 \rangle$ and $\langle R^4 \rangle$ were in agreement with known exact expressions up to the estimated statistical errors.

Fig. 1 shows a comparison of the normalized $G(r)$ from Eq. (3) to the data from our Monte Carlo simulation. Note that there is no free parameter to adjust the curves. The curves for the $d = 2$ case are qualitatively similar and the agreement between theory and the $d = 2$ Monte Carlo simulation is of equal quality. It can be improved somewhat further by applying a simple correction procedure specific to the $d = 2$ case. The general observation is that Eqs. (3), (4) and (5) reproduce the data qualitatively right and are good approximations for $t \approx 0.5$, that is $t \lesssim 2$. However, even for large values of $\kappa$ one would not expect the harmonic approximation to yield acceptable results for small $r$ since the curvature of the polymer must then be large. The somewhat surprising quality of the approximation in this region can be attributed to the fact that there the distribution function is dominated by the energy $E_{cl}(r)$ of the most probable configuration. As discussed below Eq. (3), the dominant linear term of $E_{cl}(r)$ is reproduced by the harmonic approximation, which corresponds to the well known fact that $f_r$ can be obtained by looking only at infinitesimal compressions. For $r \approx 0$, however, the deviations of $E_{cl}$ from the linear form get significant for all $\kappa$. While this is irrelevant for most applications of the distribution function due to the small probability of such configurations, it implies that the ring-closure probability $G(0)$ is not reproduced correctly by our approach. Instead one should expand the configurations around the contour of minimal energy as in Ref. [22] where the resulting expressions for $G(0)$ are evaluated numerically yielding results in agreement with our Monte Carlo data.

Another possibility to check the validity of the obtained distribution functions is to compare their moments to known results. The moments $\langle R^n \rangle$ can either be calculated from the given expressions for $G(r)$ or by applying the harmonic approximation directly to the generating function $\langle e^{fR} \rangle$. The latter method yields an expansion of $\langle R^n \rangle$ in $t$ which is seen to be correct to $O(t^2)$ when compared to the results of Ref. [17]. Calculating the moments directly from the $t = 1$ term of Eq. (4) for $\kappa(1-r) < 0.2$ and the $k = 1$ term of Eq. (5) for $\kappa(1-r) > 0.2$ produces analytical expressions which can be expanded in terms of $t$ only up to a small correction vanishing like $\exp[-\text{const.} \kappa]$. This leads to expansion coefficients that are not rational numbers but differ very slightly from the results of the generating function method (by about 0.1% for $t = 1$). Calculation from the distribution function has the advantage that averages $\langle R^n \rangle$ with arbitrary $\alpha > -d + 1$ can be obtained. The results indicate that the expansion derived in [17] for $\alpha = 2n$ with $n = 0, 1, \ldots$ is valid for all $\alpha > -d + 1$ at least up to $O(t^2)$.

A quantity of immediate experimental interest is the force-extension relation. For sufficiently stiff polymers it can be obtained from the given $G(r)$ for arbitrary forces in situations where the ends of the polymer can rotate freely. The well known strong-force limit (e.g., [8]) is reproduced by our distribution functions for arbitrary stiffness since it is determined by the behavior of $G(r)$ for $r \to 1$ where the harmonic approximation is expected to be good for all values of $\kappa$: For large forces $f$ we have $r \approx 1$ and consequently $G(r)$ will be dominated by the $t = 1$ term of Eq. (3). The relevant free energy is thus $F = -\log G(r) + f(1-r) \approx 1/(4\kappa(1-r)) + f(1-r)$, where we neglected logarithmic terms since $1 - r \to 0$ for $f \to \infty$. Since the corresponding distribution function will be strongly peaked for large $f$, $\langle r \rangle$ is given by the position of the minimum of $F$: $\langle r \rangle = 1 - 1/\sqrt{\pi f}$ which agrees with Ref. [8]. Force-extension relations for small forces can be obtained from moments of the distribution function. Our results are thus consistent with existing linear response treatments [26,27] to $O(t^2)$. If the polymer’s orientation at one end is fixed, the character of the response depends on the direction of the applied force and $G(r|u_0; L)$ is needed [24] (notation of Ref. [18]). Note that the linear response can be evaluated exactly for arbitrary values of $t$ in this case [27].

Using fluorescence microscopy, it is possible to visualize the thermal undulations of single actin filaments constrained to quasi two-dimensional configurations [13,23]. Both contour lengths and distances in embedding space can be measured from the resulting images. If locality of the interactions along the chain is assumed, different segments of one physical polymer of length $L_0$ are statistically independent. One can thus probe different length scales by obtaining experimental distribution functions $G(r; L)$ for different $L \leq L_0$. Using only $\ell_p$ as free parameter, our theoretical results can be fitted to the experimental distributions. The quality of the fits for different $L$ will indicate at what scales actin can actually be described by the wormlike chain model. This might reveal new interesting physics and help to clarify some of the ambiguities that arise in normal-mode analyses of actin flickering: The average squared amplitudes of the
normal modes with mode number \( k \) of the elastic Hamiltonian fail to decay like \( 1/k^2 \) as predicted by the wormlike chain model. Since \( G(r; L) \) can be determined without differentiation and Fourier transformation of the observed contours, the radial distribution function analysis is expected to be much more stable against unavoidable experimental errors. Another method that has been used to analyze actin flickering is the comparison of the measured decay of the tangent-tangent correlation function with the exact result \( t(s)t(s') \propto \exp(-|s-s'|/\ell_p) \). However, since an exponential decay of correlations is expected for all systems with only short range interactions, this kind of analysis is probably not very sensitive to deviations from the wormlike chain model. A very interesting possibility would be to attach two or more markers (e.g., small fluorescent beads) permanently to single strands of polymers and to observe the distribution function of the markers’ separation. This would eliminate all the experimental difficulties associated with the determination of the polymer’s contour. Note that contrary to existing methods of analysis it is not necessary to know the length of polymer between two markers; this quantity can be extracted from the observed distribution functions along with \( \ell_p \).

In conclusion, we have derived approximate analytical expressions for the end–to–end distribution function \( G(r; L) \) of semiflexible polymers. The essential ingredient in our calculation is the choice of a parameterization for the polymer’s configuration that satisfies the constraint of fixed contour length by construction. Comparison with Monte Carlo data shows good quantitative agreement for \( t = L/\ell_p \lesssim 2 \). Since the Daniels approximation is valid for \( t \gtrsim 10 \), \( G(r; L) \) of the wormlike chain is up to a crossover region now available for the whole range from rigid rods to random coils. The range of stiffness accessible to the approximations used is highly relevant for the physics of rather rigid polymers like actin. Knowledge of \( G(r; L) \) provides new possibilities for the experimental determination of \( \ell_p \) as well as the lengthscales at which real semiflexible polymers can be described by the wormlike chain model. Since the actual form of the single chain probability distribution function is an important input for the theory of many-chain systems, we hope that our work will contribute to the investigation of more complex questions such as the dynamics and viscoelasticity of networks and solutions of semiflexible polymers.

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