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Multiplectoneme phase of double-stranded DNA under torsion

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We use the wormlike chain model to study supercoiling of DNA under tension and torque. The model reproduces experimental data for a broad range of forces, salt concentrations, and contour lengths. We find a plane of first-order phase transitions ending in a smeared-out line of critical points, the multiplectoneme phase, which is characterized by a fast twist-mediated diffusion of plectonemes and a torque that rises after plectoneme formation with increasing linking number. The discovery of this phase at the same time resolves the discrepancies between existing models and experiment.

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I. INTRODUCTION

The behavior of double-stranded DNA (dsDNA) under tension and torsion plays an important role in the transcription and replication of our genetic code. The DNA present in a single human cell is long enough to outdo most of us in height; yet it is confined in nuclei with diameters in the micron range, orders of magnitude smaller than the chain would have in a θ solvent. One of the ingredients in the compactification of DNA in bacteria is supercoiling, where torsional stress results in the formation of plectonemes: loops in the molecule with the two halves of the molecule coiled around each other, like an old-fashioned telephone cord (Fig. 1). Since dsDNA forms in its relaxed state a right-handed double helix it is chiral, and a combination of torsion and tension comes automatically into play during transcription and replication. Single-molecule experiments [1] have been instrumental in investigating the elastic properties of dsDNA. The force-extension behavior of a freely rotating chain can be described by modeling the molecule as an elastic rod in a thermal environment [2,3], with all elastic strains described by just two elastic moduli: the bending modulus, and a stretch modulus S which due to its large value of ~ 700 – 1300 pN can safely be omitted for tensions in the piconewton range. This wormlike chain (WLC) model was shown [2,3] to be a good description over a large range of contour lengths and tensions.

When experimental techniques made it possible to put at the same time a torque on the molecule [4], adding the torsional degree of freedom with its linear elastic modulus to the WLC model results in a good description of the experimental data [5] for torques somewhat lower than the classical buckling transition. After this buckling transition a growing plectoneme (Fig. 1) is thought to set the slope of the force-extension curve. Many models have been constructed to predict some of the measurements but, as we will argue in this paper, are incomplete in their description of the thermal fluctuations. This led to some remarkable disagreement with experimental data. Furthermore an important feature of the phase diagram of torsionally stressed dsDNA remained uncovered. The experimental setup with which we compare the model consists of a dsDNA molecule that has one end attached to a substrate and the other end attached to a bead. This bead is either superparamagnetic (small ferromagnetic domains randomly

oriented in a polystyrene sphere) or glass. In the first case the position is controlled by the gradient of a magnetic field (hence the name magnetic tweezer), in the second by laser beams (the optical tweezer). By making use of a net magnetic moment (or a specially crafted bead in the optical case) it is possible to control the rotation of the bead. The torque is not fixed in these measurements, but the average torque is extracted either from the rotation-extension curves, or from the trap stiffness using specially crafted beads.

In this paper we will include a consistent description of thermal fluctuations, on the way adding some details to the common models that remained somewhat hidden in the usual treatments. For notational convenience we scale all energies by $k_B T$ unless explicitly mentioned. Forces will have the dimension of an inverse length and the two moduli introduced that of a length.

The setup of the paper is the following: In Sec. II we define the Hamiltonian using the contributions that are common to most models, on the way putting in some details that are not always appreciated. In Sec. III we analyze in detail the influence of thermal fluctuations on all scales. We find that in the plectoneme, the short-wavelength fluctuations renormalize the two moduli in a nontrivial way. On the global scale we analyze in depth the appearance of multiple plectonemes and find a sharp transition between multiplectoneme and single-plectoneme behavior. In Sec. IV we compare the model with several sets of experiments concerning extension and torque of the supercoiling molecule. We close the paper in Sec. V with a short discussion of some other recent models and an outlook to future developments.

II. THE HAMILTONIAN

To include a twist degree of freedom, the DNA molecule with contour length L_c is modeled as a ribbon, or equivalently a framed space curve, defined through its tangent $\mathbf{t}(s)$ and local frame rotation $\psi(s)$, with as parameter the arc length s . The number of turns Lk (linking number) we set to zero in the torsionally relaxed state. Its sign we choose positive when rotating the bead counterclockwise, as seen from the top, tightening the right-handed double helix. The gradual decrease of extension of the chain under constant tension f while increasing or decreasing Lk from zero is well described

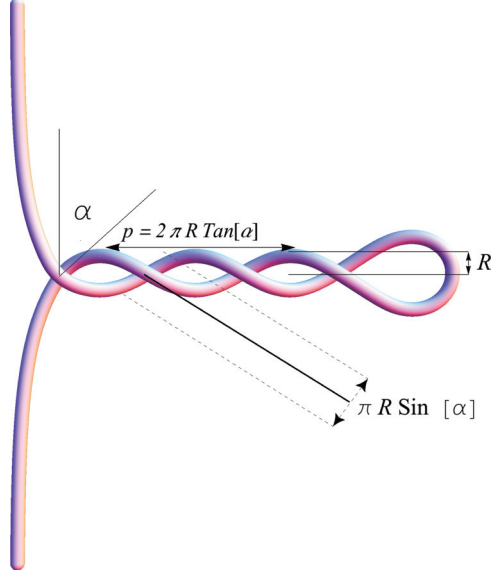


FIG. 1. (Color online) A plectoneme with plectoneme radius R , plectoneme angle α , and pitch p . The standard deviation of the fluctuation channel in the “pitch direction” is $\pi R \sin(\alpha)$.

within the framework of linear elasticity [5]. The two moduli are the (orientational) *persistence length* P_b and the *torsional persistence length* P_c . Addition of a stretch and twist-stretch coupling with a modulus that turns out to be negative [6,7], slightly improves upon this. It explains why the measured extension is not fully symmetric close to the relaxed chain. We will not consider them for the rest of the paper, since the forces below 4 pN we deal with are small compared to the experimental stretch modulus as explained in the previous section. In any case addition of a coupling term does not essentially complicate the modeling. The right handedness of the double helix on the other hand does show up in an extended plateau, already at reasonably low forces, when rotating the bead in the negative Lk direction. This is caused by the denaturation of the molecule. Since we are interested in the formation of plectonemes, we from now on restrict ourselves to the positive direction, although the same results will hold for negative Lk as long as there is no denaturation.

For the high-salt-concentration, c_s , persistence length we take $P_b(\infty) = 50$ nm, to which we add the usual electrostatic stiffening corrections following OSF theory¹ with a charge density along the chain limited by Manning condensation [10]:

$$P_b(c_s) = P_b(\infty) + \frac{1}{4\kappa^2(c_s)Q_B}, \quad (1)$$

with $\kappa(c_s)$ the inverse Debye screening length and Q_B the Bjerrum length of the solvent:

$$\kappa = \sqrt{\frac{2q_e^2 n_s}{\epsilon_r \epsilon_0 k_B T}}, \quad Q_B = \frac{q^2}{4\pi \epsilon_r \epsilon_0 k_B T}. \quad (2)$$

Here ϵ_0 is the electric constant, ϵ_r the dielectric constant of water, q_e the elementary charge, and n_s the number density

of salt molecules. For water at room temperature, the Bjerrum length is 0.715 nm. The OSF correction is small though, for example, at 20 mM and room temperature the correction term is ~ 1.6 nm.

The energy of a chain configuration up to this transition that we have to minimize has the usual elastic contributions:

$$\mathcal{E} = \int_{-L_c/2}^{L_c/2} ds \left(\frac{P_b}{2} \dot{\mathbf{t}}^2(s) + \frac{P_c}{2} \dot{\psi}^2(s) - \mathbf{f} \cdot \mathbf{t}(s) \right) - 2\pi Lk([\mathbf{t}, \dot{\psi}])\tau. \quad (3)$$

The linking number depends on the local frame rotation around the tangent, but also on the space curve the backbone traces out when traveling along the contour. The torque τ functions here as a Lagrange multiplier. The relation between Lk and the configuration of the chain we can extract from the celebrated Călugăreanu-White [11,12] relation where we imagine the chain forming part of a closed loop in which case

$$\text{Lk}([\mathbf{t}, \dot{\psi}]) = \text{Tw}([\dot{\psi}]) + \text{Wr}([\mathbf{t}]). \quad (4)$$

The twist Tw is the integrated number of turns the frame rotates around the tangent direction, Lk can be written as the Gauss integral of the two ribbon lines while the writhe (Wr) is the Gauss linking number of one of the ribbon lines with itself:

$$\begin{aligned} \text{Tw} &= \frac{1}{2\pi} \int_0^{L_c} ds \dot{\psi}(s), \\ \text{Wr} &= \frac{1}{4\pi} \oint_C \oint_C \frac{d\mathbf{r} \wedge d\mathbf{r}' \cdot (\mathbf{r} - \mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|^3}. \end{aligned} \quad (5)$$

For the fictive loop the angle function is in general multivalued, and the writhe depends on the way we close the loop. We can overcome both problems by relying on Fuller’s formula [13,14] which calculates the difference in writhe between two closed curves that are *writhe homotopic*: a homotopy of curves in the space of nonintersecting closed space curves such that nowhere along the homotopy, for any given s , is the tangent antiparallel to its value at the ends of the homotopy. In that case the writhe difference is given by [13]

$$\text{Wr}_2 - \text{Wr}_1 = \frac{1}{2\pi} \oint ds \frac{[\mathbf{t}_1(s) \times \mathbf{t}_2(s)] \cdot (\dot{\mathbf{t}}_1 + \dot{\mathbf{t}}_2)}{1 + \mathbf{t}_1(s) \cdot \mathbf{t}_2(s)}. \quad (6)$$

This formula follows from the interpretation of the writhe as the area on the direction sphere enclosed by the tangent, when going around the loop. Fuller’s formula calculates the area difference between the two homotopic curves. We consider the chain to be clamped at both ends such that the tangent and its derivative are fixed at the ends. Defining the zero of the chain’s writhe to be the torsionally relaxed state, we can calculate the writhe of any writhe homotopic perturbation under the clamped boundary conditions from Eq. (6) integrated over the chain, effectively keeping the closing part invariant. Implicitly we assume the bead is large enough, compared to chain fluctuations, that the chain will not change linking number by looping over the sphere.

We choose the twist angle coordinate to be zero at the substrate. A linear stability analysis around the straight configuration is now straightforward showing that there is a

¹OSF stands for Odijk [8] and Skolnick, Fixman [9].

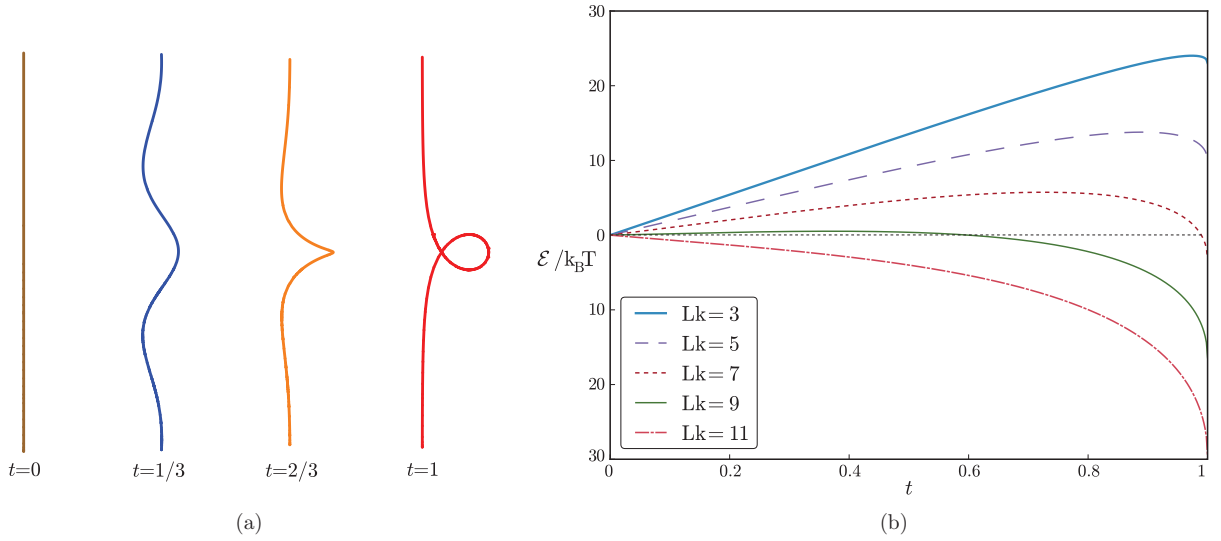


FIG. 2. (Color online) The homoclinic solutions. (a) Curves of the homoclinic solutions for the homoclinic parameter $t = 0, 1/3, 2/3$, and 1. (b) The energy of the homoclinic solutions relative to the straight rod energy, where all the linking number is in the twist of the chain. Here $L_c = 600$ nm and $f = 2$ pN $\gg f_0$.

bifurcation point at

$$Lk_{cr} = L_c \frac{\sqrt{f P_b}}{\pi P_c}. \quad (7)$$

Before reaching this bifurcation point other local minima start to appear, which have to be taken into account in a thermal environment. The energy minima of the Hamiltonian that we are looking for should fulfill the boundary conditions of clamped ends with tangents parallel to the tension. The homoclinic solutions of an elastic rod under tension fulfill these boundary conditions in the infinite-rod limit. They form a one-parameter family of localized helices [see Fig. 2(a)], ranging from the straight rod to a localized loop in spherical tangent coordinates given by [15]

$$\cos \theta(s, t) = 1 - 2t^2 \text{sech}^2 \left(\frac{st}{\lambda} \right), \quad (8)$$

$$\phi(s, t) = \arctan \left[\frac{t}{\sqrt{1-t^2}} \tanh \left(\frac{st}{\lambda} \right) \right] + \sqrt{1-t^2} \frac{s}{\lambda},$$

with $t \in [0, 1]$ and $\lambda = \sqrt{P_b/f}$ the deflection length, the length scale above which the tension dominates thermal fluctuations. Each of these solutions is valid for a specific torque and is not a ground state in the supercoiling problem. They do function though as lowest col over the barrier towards the almost closed loop that forms the start of a plectoneme. This can be shown in a straightforward manner starting from Eq. (3), using spherical coordinates for the tangent field. The twist term we can drop for the analysis. The writhe of the chain is a continuous map of the space curves that form the homotopy connecting the straight curve and the almost closed loop. The Euler-Lagrange equations are easy to solve using the boundary conditions $\theta(\pm L_c/2) = 0$ and solving τ for a fixed writhe, we find that the homoclinic solutions are indeed the extrema of the solutions with this writhe. The second functional derivative shows them to be minima.

When traversing the homoclinic solutions from $t = 0$ to $t = 1$, the writhe and bending energy of the chain are given by

$$W_{r_{loop}}(t) \simeq \frac{1}{2\pi} \int_{-\infty}^{\infty} ds \frac{\hat{e}_z \times \mathbf{t}(s, t) \cdot \dot{\mathbf{t}}(s, t)}{1 + \hat{e}_z \cdot \mathbf{t}(s, t)} = \frac{2}{\pi} \arcsin(t), \quad (9)$$

$$\mathcal{E}_{loop} = 2f L_{loop} = 8f \lambda t,$$

with L_{loop} denoting the decrease in extension of the chain which we identify with the loop length. When Lk is kept constant, the increasing writhe decreases the twist following Eq. (4), resulting in a loop energy at constant Lk of

$$\mathcal{E}(t) = \mathcal{E}_{loop} + \frac{2\pi^2 P_c}{L_c} \left(Lk - \frac{2}{\pi} \arcsin(t) \right)^2. \quad (10)$$

From this expression it follows that for tensions $f < f_0 := 4P_c^2/(P_b L_c^2)$ the energy minimum shifts from the straight rod continuously to the homoclinic loop when the linking number is increased from Lk_{cr} (7) till 1. For tensions above f_0 only a limited range of stable solutions in between the two extrema exists. Also in that case the straight rod ceases to be stable at Lk_{cr} , while the barrier to the loop solution disappears a little later when

$$Lk = Lk_{cr} \sqrt{1 - \frac{4}{Lk_{cr}^2 \pi^2}} + \frac{2}{\pi} \arcsin \left(\frac{2}{Lk_{cr} \pi} \right). \quad (11)$$

In Fig. 2(b) a typical situation is sketched for a chain of 600 nm and a tensile force of 2 pN $\gg f_0$. Note how already in an early stage a local minimum starts to form separated from the straight rod by a barrier and how that barrier moves to smaller t values with increasing Lk .

When t approaches 1, the closed loop, excluded volume interactions have to be taken into account. DNA is, at neutral pH, a strong polyelectrolyte with one charge per backbone phosphate. In a thermal environment the interaction between

two chains approaching each other at a large angle is a steep potential, at a distance not far from the Debye screening length [16]. A point of closest approach exists in homoclinic solutions whenever $t > t_c \simeq 0.804\,24$. Its value, $d_{\min}(t)$, is the nontrivial minimum of

$$d(s, t) = 2\lambda \sqrt{4t^2 \operatorname{sech}^2 \frac{st}{\lambda} \sin^2 \frac{s\sqrt{1-t^2}}{\lambda} + \left(\frac{s}{\lambda} - 2t \tanh \frac{st}{\lambda}\right)^2}. \quad (12)$$

Within the range $t \in [t_c, 1[$ we can approximate d_{\min} with

$$d_{\min}(t) = 2\lambda \left(\sqrt{\frac{1-t}{0.3799}} - 0.001\,12 \right). \quad (13)$$

For a given force this distance has a maximum of $d_{\min}(t_c) \simeq 1.4\lambda$. The point of closest approach functions as a pivot point from which the plectoneme nucleates as long as it is energetically cheaper to reduce the twist through a writhing plectoneme than through the writhe of another homoclinic loop.

The radius R and angle α of the plectoneme are set by a delicate balancing of a variety of contributions. The electrostatic repulsion, the bending energy, and the entropic repulsion all depend on and directly influence R and α . Indirectly they, like the tension, influence the parameters through the writhe efficiency of the plectoneme. This forms the basis for most of the modeling done for plectoneme formation. For the electrostatic repulsion we use the results from Ref. [17]:

$$\begin{aligned} \epsilon_{\text{el}}^0(R, \alpha) &= \frac{q_{\text{eff}}^2 Q_B}{2} \sqrt{\frac{\pi}{\kappa R}} e^{-2\kappa R} Z[\cot(\alpha)], \\ Z(x) &= 1 + m_1 x^2 + m_2 x^4, \\ m_1 &= 0.828, \quad m_2 = 0.864, \end{aligned} \quad (14)$$

valid for $\cot(\alpha) < 1$, with q_{eff} the effective charge density of the centerline of a cylinder which is the source of a Debye-Hückel potential that coincides asymptotically, in the far field, with the nonlinear Poisson-Boltzmann potential of that cylinder with a given surface charge. For dsDNA we take a naked charge density of two charges per 0.34 nm, representing the two phosphate charges per base pair, and a radius of 1.0 nm. The expansion is a fit that behaves reasonably also for $\cot \alpha$ close to 1, where a standard asymptotic expansion would fail. The effective charge density q_{eff} is finally calculated following Ref. [18].

In contrast to the persistence length corrections, these calculations are based on the bare charge of the DNA chain. It can be shown that Manning condensation follows asymptotically [19]. Note that by using an effective potential based on the Poisson-Boltzmann equation the model already includes thermal motion of counterions and salt ions. We will nonetheless refer to the model in this section as being athermal.

Note further that we use the usual simplification of taking the plectoneme radius and angle to be constant along the plectoneme. We set the homoclinic parameter t by the demand that the nontrivial shortest distance between the two legs of the homoclinic solution equals twice the plectoneme radius. It is here that we will define the start of the plectoneme. The

remaining part of the homoclinic solution stays connected to the end of the plectoneme and rotates around the plectoneme axis with growing plectoneme length. In this way our solution is continuous, although not in general differentiable. One could argue that the assumption of constant plectoneme parameters does not represent the true minimum of the free energy and that in reality the space curve should be smooth. However these are details of the energetics that are not important for the experiments, where most contributions come from the plectoneme alone. The plectoneme has, in addition to the potential energy density, caused by the tension, the usual energy density contributions of bending:

$$\epsilon_{\text{bend}}(R, \alpha) = \frac{P_b \cos^4(\alpha)}{2} \frac{1}{R^2}. \quad (15)$$

The writhe density of the plectoneme is given by the well-known expression

$$\omega = \frac{\sin(2\alpha)}{4\pi R}. \quad (16)$$

This expression is often not appreciated. The naive approach of calculating the writhe density using Fuller's equation relative to the plectoneme axis does result, upon averaging, in the right expression but neglects the influence of the end loop and is relative to the wrong axes. Arguing that the end loop is only a short stretch of the chain and thus negligible is clearly wrong since every turn of the plectoneme length contributes equally to the writhe of the end loop. In Appendix A it is shown that in fact expression (16) is right compared to the tension axes only when the end-loop contribution to the plectoneme writhe is included.

Putting the ingredients together we find for the energy of the chain with plectoneme

$$\begin{aligned} \mathcal{E}(R, \alpha) &= \mathcal{E}_{\text{loop}}(t(R)) + L_p \epsilon_{\text{plect}}(R, \alpha) \\ &\quad + \frac{2\pi^2 P_c}{L_c} (\text{Lk} - W_{\text{rloop}} - L_p \omega)^2, \\ \epsilon_{\text{plect}}(R, \alpha) &:= f + \epsilon_{\text{bend}}(R, \alpha) + \epsilon_{\text{el}}^0(R, \alpha). \end{aligned} \quad (17)$$

The plectoneme contour length L_p is found by minimizing the energy:

$$L_p = \frac{\text{Lk} - W_{\text{rloop}}}{\omega} - \frac{\epsilon_{\text{plect}} L_c}{4\pi^2 P_c \omega^2}, \quad (18)$$

where we assumed Lk to be large enough to make L_p positive. For a long enough chain and plectoneme the loop contribution can be neglected in determining the optimal values for the plectoneme parameters R and α . This infinite-chain limit is the usual approach in modeling the plectoneme and is already implicitly included in the electrostatic contribution (14). The price we pay for this simplification is small, at most noticeable close to the transition.

Starting from the torsional relaxed chain, after introducing a certain number of turns lower than the critical linking number, a solution containing a plectoneme will appear with an energy equal to the straight solution. More precisely, the buckled configuration at the transition either has a finite plectoneme that minimizes the energy or consists of only

the loop:

$$L_{p,tr} = \begin{cases} 0 & \text{in case } \Delta \leq 0 \text{ or} \\ & \sqrt{\frac{L_c \Delta}{2\pi^2 P_c}} < W r_{loop}, \\ \left(\sqrt{\frac{L_c \Delta}{2\pi^2 P_c}} - W r_{loop} \right) \frac{1}{\omega} & \text{otherwise,} \end{cases} \quad (19)$$

$$Lk_{tr} = \begin{cases} \frac{\epsilon_{eplect}}{4\pi^2 \omega} \frac{L_c}{P_c} + \sqrt{\frac{L_c \Delta}{2\pi^2 P_c}} & \text{if } L_{p,tr} > 0, \\ \frac{\epsilon_{loop}}{4\pi^2 W r_{loop}} \frac{L_c}{P_c} + \frac{1}{2} W r_{loop} & \text{if } L_{p,tr} = 0, \end{cases}$$

with

$$\Delta := W r_{loop} \left(\frac{\epsilon_{loop}}{W r_{loop}} - \frac{\epsilon_{eplect}}{\omega} \right) \quad (20)$$

the cost per writhe difference between loop and plectoneme. In this nonthermal model plectoneme formation will not happen when $\Delta < 0$ since it is always cheaper to form a new loop than to grow a plectoneme, but entropic contributions that we will treat in the next section will change that. This transition point is marked by a drop in extension that is partly due to the homoclinic loop, partly due to the length of the plectoneme at the transition. Although this transition is not sharp a local minimum leading to the plectoneme does not appear until Lk has reached a value Lk^0 where either

(1) the plectoneme length minimizing the energy (18) has reached zero: $Lk_0 = \frac{\epsilon_{eplect} L_c}{4\pi^2 P_c \omega} + W r_{loop}$, or

(2) the homoclinic solution has reached the maximum of the energy barrier at t_R and marking the formation of a local minimum at zero plectoneme-length: $Lk^0 = L_c \frac{P_b \sqrt{1-t_R}}{P_c \pi \lambda} + \frac{2}{\pi} \arcsin(t)$.

In any case we see that in the infinite-chain limit Lk^0 scales as Lk_{tr} with the contour length. Therefore we will in the following switch to linking number densities $lk := Lk/L_c$.

The plectoneme length depends on both tension and salt concentration, but on top of that scales with the square root of the contour length. This has some interesting consequences when considering the appearance of multiple plectonemes. In case the ground state at the transition has a finite-size plectoneme length, the number of plectonemes in general does not grow with the system size. This in contrast with a situation where $\Delta \leq 0$. This will become a point size defect in the infinite-chain limit and results in a finite density of plectonemes. Roughly speaking, increasing f or decreasing the salt concentration c_s increases the energy per writhe of the plectoneme, thereby shortening its start length. This leads to the following picture in the f, c_s, lk space: for high c_s and low f there is a first-order-like transition from the plectonemeless configuration to a finite-length plectoneme. The jump in extension scales with the square root of the chain length. These transition points are like a plane of first-order transitions dominated by the finite length of the starting plectoneme. The plane ends in a line of continuous transitions where the transition is from straight to a configuration with an increasing number of plectonemes resulting in a finite plectoneme density: the *multiplectoneme* phase. A drop in extension caused by the end loop can still be present for short chains but thermal fluctuations smooth the transition for longer chains.

This “multiplectoneme phase” has some interesting, biologically relevant, dynamical properties that we will come back to in the next section.

III. THERMAL FLUCTUATIONS AND THE MULTIPLECTONEME PHASE

To account for thermal fluctuations several strategies have been employed in modeling plectoneme formation. The simplest strategy is to ignore them [20–22] at most adding an overall chain shortening factor [22] that does not change the slope. Another strategy is to ignore only thermal fluctuations in the plectoneme [23,24], arguing that at least for higher tensions the fluctuations are small and can as a consequence be neglected. To account for the entropic repulsion of the strands’ confinement an entropic term from older bacterial supercoiling models is added as an independent ingredient [24,25].

In the first case, it is not clear why the size of the thermal fluctuations inside the plectoneme should be the same as in the tails. The confinement of the chain in the plectoneme is the result of a subtle equilibrium between the applied tension, the electrostatic repulsion, and the need to reduce the twist through writhe. Furthermore this procedure needs an extra surface charge reduction of the chain to reproduce experimental slopes [22].

The second approach (fluctuations in the plectoneme are small), when properly applied, does not need this charge reduction to get a reasonable agreement with some of the experiments (as long as the salt concentration is not too low) but has the conceptual problem that there is no *a priori* reason why the plectoneme would be totally immune to fluctuations. The reasoning that thermal fluctuations are small within the plectoneme and thus can be ignored is erroneous since the plectoneme free energy has to be compared with the tails where the finite fluctuations have a known dependence on tension and applied torque. The only conclusion one can draw, following this line of thought, is that the extreme reduction in the number of configurations prohibits plectoneme formation.

The last approach ignores the influence of torsion although this torsion is strongly influencing the free energy in the tails. Furthermore the bending energy density and the writhe density of the plectoneme are both affected by thermal fluctuations. In the following we will model thermal fluctuations in the plectoneme with the same rigor as was done previously [5] for the tails.

A. Short-wavelength fluctuations

Below the transition we use the results from Moroz and Nelson [5]. These can be extended [26] with a finite stretch modulus $S \simeq 300 \text{ nm}^{-1}$ [25] and twist-stretch coupling $B \simeq -21$ [25]. Including these moduli affects the (reciprocal) expansion parameter K as introduced in Ref. [5]:

$$K = \sqrt{f P_b - \left(\pi P_c' lk + \frac{Bf}{2S} \right)^2} \quad (21)$$

with $P_c' := P_c - B^2/S$ the effective torsional persistence length from Ref. [26]. The free energy density of the chain

expressed in this factor can then be written as [5,26]

$$\begin{aligned} f_{\text{tail}} &= 2\pi^2 P_c \text{lk}^2 - \frac{(f - 2\pi B \text{lk})^2}{2S} \\ &\quad - f + \frac{K}{P_b} \left(1 - \frac{1}{4K} - \frac{1}{64K^2} \right) \\ &\simeq f_{\text{tw}}^t - f \left(1 + \frac{f - 4\pi B \text{lk}}{2S} \right) + \frac{1}{\lambda} \left(1 - \frac{\lambda}{4P_b} - \frac{\lambda^2}{64P_b^2} \right), \end{aligned} \quad (22)$$

with the twist free energy density

$$f_{\text{tw}}^t \simeq 2\pi^2 P_c' \langle tw^2 \rangle = 2\pi^2 P_c' \left(1 - \frac{\lambda P_c'}{4P_b^2} \right) \text{lk}^2. \quad (24)$$

Since the maximum tensions applied stay below 1 nm^{-1} (4 pN) the effect of these moduli stays small thanks to the relatively strong resistance against stretching, and we will drop them in the rest of the paper, by setting $S = \infty$, to decrease the clutter.

The twist energy is one of the main results of Moroz and Nelson [5] who introduced the notion of a *thermally* renormalized torsional persistence length:

$$P_c^{\text{ren}}(\lambda) = \left(1 - \frac{\lambda P_c}{4P_b^2} \right) P_c. \quad (25)$$

The linking number that was put into the chain gets spread between twist and a thermal writhe that is not symmetric around the straight twisted rod but has a directionality, thereby decreasing the twist density and apparently decreasing P_c . The expectation value of this thermal writhe density, $\omega_{\text{tail}}^{\text{th}}$, and the resulting thermal shortening, ρ_{tail} , both up to lowest order, are given by

$$\begin{aligned} \langle \omega_{\text{tail}}^{\text{th}} \rangle &= \frac{P_c \lambda}{4P_b^2} \text{lk}, \\ \rho_{\text{tail}} &= 1 - \frac{1}{2K} \left(1 + \frac{1}{64K^2} + \dots \right) \\ &\quad + \left(\frac{1 - \coth\left(\frac{L_c K}{P_b}\right)}{2K} + \frac{P_b}{2L_c K^2} \right). \end{aligned} \quad (26)$$

The last term in Eq. (27) is a finite-size correction, which we will also drop in the following.

The validity of these expressions is limited to values of force and linking number that make the expansion factor K large enough. Moroz and Nelson argued that for $K^2 > 3$, the error in the extension should be below 10%, based on a comparison with the next term in the asymptotic expansion.

There are in fact two other sources for errors: the appearance of knotted configurations, which should have been excluded from the partition sum, and configurations with a writhe that differs by a multiple of 2 from the calculated writhe caused by the use of Fuller's equation. For large K when large deviations from the straight rod are highly suppressed the influence of these effects is small, and we will consider a value of $K^2 = 3$ to be the lower bound below which the theoretical treatment of Ref. [5] breaks down.

Once a plectoneme is formed we can think of three distinct regions: the tails, which can be treated as the straight solution, the end loop, and the plectoneme.

As shown in Ref. [27], in a WLC under tension, the length of a loop, not the contour length of the chain forming the loop, is to lowest order unaffected by thermal fluctuations. This was shown for a loop with homoclinic parameter $t = 1$ with the two tails bound by a gliding ring at the contact point. There is no reason to doubt that this will hold also for the end loops of the plectonemes, since they are sufficiently close to the closed loop, with the essential difference that the tails are not bound together but lie in an effective potential well resulting from a twist-induced attraction and an electrostatic repulsion. Thermal fluctuations necessarily open the loop from its ground state value, thus decreasing its length. This loop destabilization effect becomes unimportant for a finite-size plectoneme configuration, since loop opening and plectoneme radius are linked. To avoid unnecessary complications we will just ignore the entropic loop contributions and instead determine the relevant loop size from the plectoneme. It is possible to add electrostatic interactions to the loop [28], but the advantage of not having to estimate these and the contribution of the entropic repulsion to the end-loop free energy more than compensates for the small error it might produce in the free energy close to a possible plectonemeless loop configuration. In general this simplification hardly affects the jump in length seen in the turn extension plots at the transition, since jumps indicate usually a finite-size plectoneme at the transition, while the plectoneme parameter has only a limited range in light of the lower limit t_c .

The plectoneme part needs a more careful examination. We start from the calculations from Ref. [17]. They considered one strand of the regular plectoneme fluctuating in the mean field potential of the opposing strand, assuming the fluctuations to have a Gaussian distribution around their average in two directions perpendicular to the strand. One direction is chosen pointing towards the opposing strand, the radial direction, and the other normal to this direction, the pitch direction. Fluctuations in the radial direction are dominated by the exponent of the electrostatic interactions, while fluctuations in the pitch direction have much less influence on the energetics. We stress the advantage of this approach over the expansion of the effective confining potential around the ground state. In the radial direction the potential is highly skewed, exponentially increasing towards smaller radius. A harmonic approximation would be valid only in a tiny region around the ground state. Instead we assume fluctuations small compared to its typical length scale, the persistence length. Denoting the standard deviation of the Gaussian distribution in the radial and pitch direction by respectively σ_r and σ_p , the electrostatic part of the free energy changes approximately to [17]

$$\begin{aligned} f_{\text{el}}(t, \alpha, \sigma_r) &= \epsilon_{\text{el}}^0 e^{4\kappa^2 \sigma_r^2} \\ &= \frac{q_{\text{eff}}^2 Q_B}{2} \sqrt{\frac{\pi}{\kappa R(t)}} e^{4\kappa^2 \sigma_r^2 - 2\kappa R(t)} Z[\cot(\alpha)]. \end{aligned} \quad (28)$$

The steep exponential rise of this free energy contribution clearly limits the value of σ_r to be of order $(2\kappa)^{-1}$. This distinguishes the magnitude of radial fluctuations from those in the pitch direction.

It has been argued [29] that the standard deviation in the pitch direction should be of the order of the pitch itself. This result one expects also on geometrical grounds, as shown

in Fig. 1. While an exact value is hard to obtain, it is considerably larger than σ_r . As it is the tightest direction that dominates the free energy of confinement [30], our results are fairly insensitive to its precise value. In the following we chose $\sigma_p = \pi R \sin(\alpha)$, which is the standard deviation of the channel formed by the two neighboring stretches of fluctuating opposing strand. The undulating chain contracts with a factor ρ_{pl} , that we will discuss further below. This contraction on the other hand decreases the bending energy density and the writhe density of the plectoneme in a nontrivial way. In Appendix B it is shown that they change to

$$\begin{aligned} \epsilon_{\text{bend}} &\rightarrow f_{\text{bend}} = \rho_{\text{pl}}^4 \epsilon_{\text{bend}} = \rho_{\text{pl}}^4 \frac{P_b \cos^4(\alpha)}{2 R^2}, \\ \omega &\rightarrow \rho_{\text{pl}} \omega = \rho_{\text{pl}} \frac{\sin(2\alpha)}{4\pi R}. \end{aligned} \quad (29)$$

To compute the entropic cost of confinement, we cannot neglect the twist in the chain. The twist along the backbone couples to the other degrees of freedom mostly through the global constraint encoded in White's equation (4). As one expects, and was experimentally shown [31], twist relaxation is fast compared to tangential fluctuations. This allows us to integrate out these fast modes and take the twist free energy density to be constant throughout the chain.

In the tails thermal writhe is suppressed by the tension, while in the plectoneme it is suppressed by the confinement caused by a combination of electrostatics, tension bending, and twist. These thermal writhes are in general not the same even when their twist energy densities are. Therefore we need to take the thermal writhe in the plectoneme explicitly into consideration. We assign part of the total linking number to the tails and loop, from which follows a tension-dependent expectation value of thermal writhe and twist density according to Eq. (26). The rest of the linking number has to be accounted for by the plectoneme. We use this difference as the definition of its linking number. A large part of this linking number is stored in the twist and writhe of the zero-temperature plectoneme, but partly it exists in the thermal writhe of the strands of the plectoneme. For the calculation of the relevant quantities of a torsionally constrained confined WLC, we assume we can capture the physics of confinement of the plectoneme strands with that of a chain confined by a harmonic potential with the same standard deviations σ_r and σ_p . In other words, the transverse distribution is Gaussian enough. The relevant calculations for the confinement problem were performed in Ref. [30]. The free energy density of a confined WLC as a function of linking number density lk_{str} and the standard deviations in the two orthogonal channel directions σ_r and σ_p is to lowest order

$$\begin{aligned} f_{\text{strand}} &= f_{\text{tw}}^{\text{str}} + \frac{3}{8} \left(\frac{1}{\lambda_r} + \frac{1}{\lambda_p} \right), \\ \text{with } f_{\text{tw}}^{\text{str}} &:= 2\pi^2 P_c \langle \text{tr}_{\text{str}}^2 \rangle = 2\pi^2 P_c^{\text{ren}}(\lambda_s(\sigma_r, \sigma_p)) \text{lk}_{\text{str}}^2, \end{aligned} \quad (30)$$

where $P_c^{\text{ren}}()$ is the same function of λ as given by Eq. (25). The effective deflection length λ_s , the length scale over which the confining potential starts to dominate thermal fluctuations,

is given by [30]

$$\lambda_s = 2 \frac{\lambda_r^3 \lambda_p + \lambda_r^2 \lambda_p^2 + \lambda_r \lambda_p^3}{(\lambda_r + \lambda_p)(\lambda_r^2 + \lambda_p^2)}, \quad \lambda_{r,p} := (P_b \sigma_{r,p}^2)^{1/3}. \quad (31)$$

The first term of Eq. (30) is the twist free energy density, the second term is the entropic cost of confinement. Note that the confining potential, due to bending and electrostatics, is *not* included [17,30]. To the same order, the contraction of the polymer is found to be

$$\rho_{\text{pl}} = 1 - \frac{1}{4} \left[\frac{\lambda_r}{P_b} + \frac{\lambda_p}{P_b} \right], \quad (32)$$

which is up to this order equal to the torsionless contraction; inclusion of stretch and stretch-twist moduli or higher-order terms changes this. From Eq. (31) we see that in the case $\sigma_r \ll \sigma_p$ the effective deflection length is reduced to $\lambda_s \simeq 2\lambda_r$, and indeed it is the tightest direction that sets the free energy as alluded before.

These results are valid for undulations in, and thermal writhe with respect to, a straight channel. However the writhe, as a local observable, is defined only with respect to a reference curve, which is the writhing plectoneme. In Appendix B it is shown that, under reasonable assumptions, thermal writhe can be treated as an additive correction to the plectoneme writhe, where the thermal writhe is calculated as the thermal writhe of an undulating chain with a finite linking number, confined to a straight channel.

The reason is that the length scale over which the fluctuation channel axis can be considered straight is of the order of the contour length over which the r and p directions rotate around the channel axis which is of the order of the pitch or, as argued above, the standard deviation in the pitch direction. In all relevant cases the standard deviation in the radial direction is considerably smaller than in the pitch direction. Since it is this length scale, associated with the tightest direction, that determines the influence of confinement on the free energy, the energetics of the global writhing path decouples from the thermal fluctuations. The contraction ρ_{pl} depends on fluctuations in the pitch direction and therefore its size does affect plectoneme formation. The free energy density of the plectoneme is the sum of this confinement, the bending (29), and electrostatic (28) free energy:

$$f_{\text{plect}} = f_{\text{bend}} + f_{\text{strand}} + f_{\text{el}}. \quad (33)$$

Equating f_{tw}^{t} and $f_{\text{tw}}^{\text{str}}$ allows us to eliminate the linking density of the strands in the plectoneme as a parameter and write $\text{lk}_{\text{str}} = (1 - \epsilon) \text{lk}$, with

$$\epsilon = 1 - \sqrt{\frac{P_c^{\text{ren}}(\lambda)}{P_c^{\text{ren}}(\lambda_s)}} \quad (34)$$

small but in general nonzero. This is indeed the case in all experimental conditions studied: For forces ranging from 0.5 to 4 pN and salt concentrations from 20 to 320 mM, a crude estimate is easily made, namely, $\epsilon \in [0, 0.1]$. Although the difference in “thermal waste” while transforming linking number into twist is rather small, it would be wrong to draw the conclusion that entropic effects can be neglected, since the entropic part of the free energy varies as $\simeq k_B T / \lambda$. The difference between the two states can be up to $1 k_B T / \text{nm}$.

It is worthwhile to split off the twist contribution to the free energy densities:

$$\left. \begin{matrix} f_{\text{tail}} \\ f_{\text{plect}} \end{matrix} \right\} = f_{\text{tw}} + \left\{ \begin{matrix} g_{\text{tail}} \\ g_{\text{plect}} \end{matrix} \right. \quad (35a)$$

with

$$\begin{aligned} g_{\text{tail}} &= -f + \frac{1}{\lambda} \left(1 - \frac{\lambda}{4P_b} - \frac{\lambda^2}{64P_b^2} \right), \\ g_{\text{plect}} &= \frac{3}{8} \left(\frac{1}{\lambda_r} + \frac{1}{\lambda_p} \right) + f_{\text{bend}} + f_{\text{el}} \end{aligned} \quad (35b)$$

the remaining free energy contributions. We will use $\Delta g = g_{\text{plect}} - g_{\text{tail}}$ to denote their difference. Once a plectoneme has formed, the expectation value of its contour length follows from the combined linking numbers of plectoneme and end loop, which should add to the linking number that was externally applied:

$$\begin{aligned} \text{Lk} &= (L_c - L_p)\text{lk} + L_p[\rho_{\text{pl}}\omega + (1 - \epsilon)\text{lk}] + W_{\text{rloop}} \\ \Rightarrow l_p &:= \frac{L_p}{L_c} = \frac{\nu - \text{lk} - W_{\text{rloop}}/L_c}{\rho_{\text{pl}}\omega - \epsilon\text{lk}}, \end{aligned} \quad (36)$$

with $\nu := \text{Lk}/L_c$ the applied linking number density. The reduced free energy density of this one-plectoneme configuration

and its extension are

$$\begin{aligned} f_1 &= (1 - l_p)f_{\text{tail}} + l_pf_{\text{plect}} \\ &= f_{\text{tw}} + g_{\text{tail}} + l_p\Delta g + \frac{\mathcal{E}_{\text{loop}}(t)}{L_c}, \end{aligned} \quad (37)$$

$$\bar{z} := \frac{z}{L_c} = \rho_{\text{tail}}(1 - l_p) - \frac{L_{\text{loop}}}{L_c}, \quad (38)$$

both depending on the four parameters R (or t), σ_r , α , and lk . The calculation comes down to a four-parameter minimization procedure. The resulting plectoneme angle is almost independent of applied tension or salt concentration; see Fig. 3(b). This is a result of the $Z(\alpha)$ term in Eq. (14), reflecting the influence of the electrostatic repulsion to counter the demand for writhe efficiency (low α). Using this concept of energy per writhe gained in the plectoneme also helps in understanding the general trend of the plectoneme radius as shown in Fig. 3(a). Increasing the tension decreases the radius to counter the growing energy per writhe. The same holds for an increase of the range of the electrostatic repulsion, by lowering the salt concentration. Note that the plectoneme radius is always large enough for the (reduced) electrostatic potential to be below 1 in the overlap region in between the two strands. This is needed to justify the use of the Debye-Hückel tails in calculating the effective potential between the strands [32].

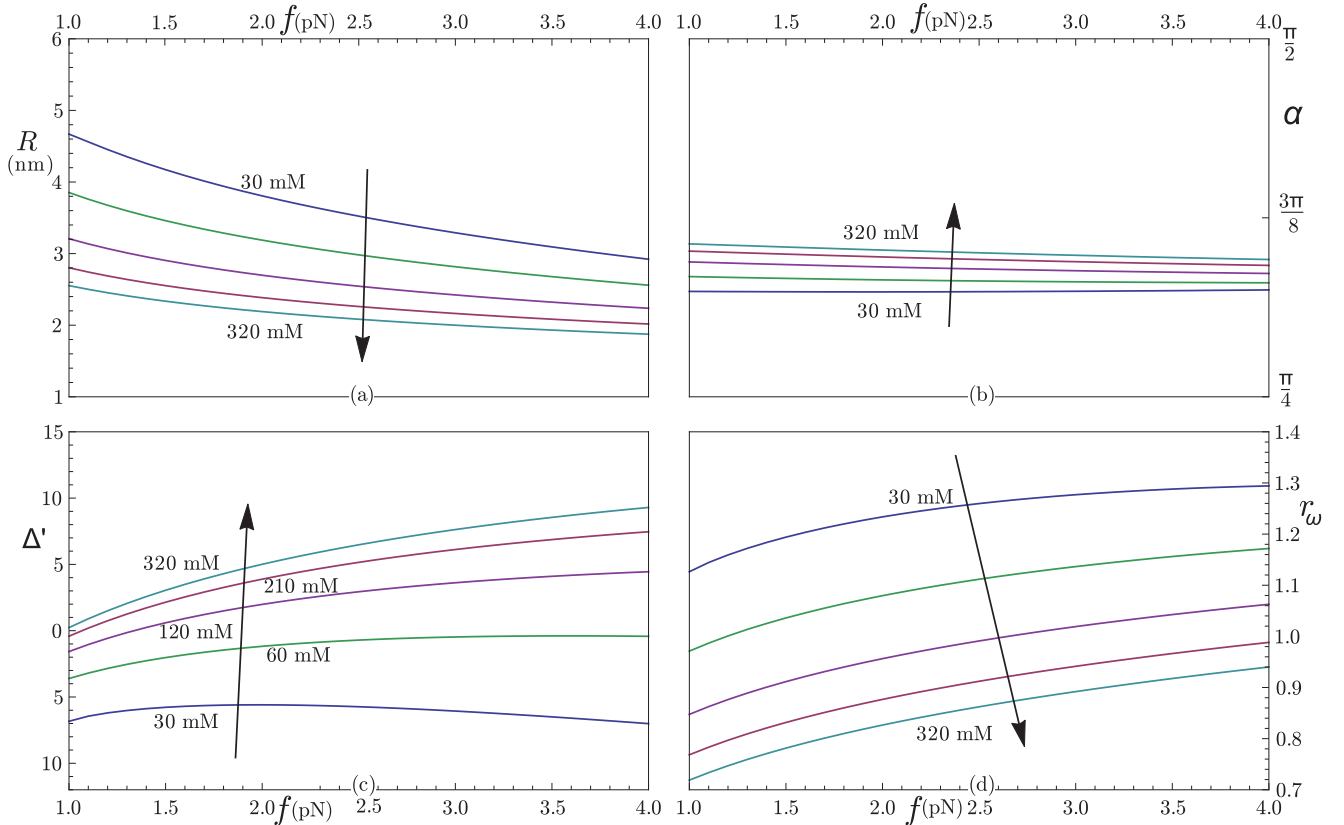


FIG. 3. (Color online) Force dependence of (a) the plectoneme radius, (b) the plectoneme angle, (c) the energy per writhe difference between loop and plectoneme with logarithmic correction related to the choice of cutoff, and (d) the writhe density ratio between loop and plectoneme for salt concentrations of 30, 60, 120, 210, and 320 mM. The arrows point in the direction of increasing salt. The range for α was on purpose chosen to be the full allowed range for a stable plectoneme, showing that its value is hardly dependent on the environment.

In the long-chain limit with finite plectoneme length the loop contribution can be neglected in determining the four parameters. We can assume that ϵ is small compared to ω , under conditions where a plectoneme forms. We can also neglect the dependence of ρ_{pl} on the parameters; its variational contribution is on the order of $\lambda_{r,p}/P_b$, which is small by assumption. The long-chain finite-plectoneme free energy is

$$f_1 = f_{\text{tw}}(\text{lk}) + g_{\text{tail}} + \frac{\nu - \text{lk}}{\rho_{\text{pl}}\omega(R, \alpha)} \Delta g(R, \alpha, \sigma_r). \quad (39)$$

The linking number density and chain extension are readily obtained in this limit:

$$\text{lk} = \frac{\Delta g}{4\pi^2 P_c \rho_{\text{pl}}\omega(R, \alpha)}, \quad \bar{z} = -\frac{\rho_{\text{tail}}}{\rho_{\text{pl}}\omega(R, \alpha)}. \quad (40)$$

Minimizing the free energy within this approximation is equivalent to minimizing the linking number density. This is not really a surprise since plectoneme formation is driven by linking number.

A numerical minimization gives results that compare reasonably well with experiments. The transition point and height of the jump at the transition as well as the slope after the transition are within experimental error for high enough forces and salt concentrations; see the dotted lines in Fig. 7. The lack of agreement at low salt clearly inversely correlates with K^2 . Dropping the assumption of equal linking number densities in tail and plectoneme hardly improves the results, even when the value of K^2 stays well above 3. This discrepancy, which is slightly stronger when fluctuations are neglected, has led to a variety of speculations, such as an effective charge reduction [22], or a charge correlation effect between the two intertwined superhelices that form the plectoneme [24]. The deviation of the experimental slopes from the calculated one goes hand in hand with the decrease of the height of the potential barrier between straight and plectoneme configurations. But our theory is not complete yet: the inclusion of other local minima in addition to these two configurations turns out to be of greater importance than has been acknowledged until now, as we will show in the next section.

B. Tunneling to the plectoneme

Contributions of local minima have to be taken separately into account in any perturbative calculation. Accepting the simplification that a plectoneme has a well-defined radius and angle that are length independent, the only concern is the barrier height between $t = 0$ and its final value t_R corresponding to the plectoneme radius.

The usual way to take these local minima into account is to treat them as a gas of defects that compete with their entropic gain against the energetic advantage of the ground state. This is the situation that would exist in a torque-regulated setup. In our case where the linking number is the control parameter the treatment changes essentially. A defect changes the linking number and so the energy of the configuration in which it is embedded. Furthermore the defects are themselves plectonemes and so to understand thermal fluctuations close to the transition we actually study multiple plectoneme configurations. Multiple plectonemes were

considered before [7,33] but mostly seen as small corrections on the one-plectoneme configurations.

The entropic gain of a multiple plectoneme configuration is twofold: there is the usual combinatoric positional freedom of defect placement (the “gas of defects”), but there is also an increase in configurations due to the freedom in distributing the total plectoneme length over the individual plectonemes. Treating the plectonemes as having a hard-core repulsion, one finds for the partition sum of a configuration with total plectoneme contour length $L_p(m)$ spread out over m plectonemes

$$Z_m = \frac{\sqrt{L_c}}{\Lambda^{2m-3/2}} \frac{L_p^{m-1}(m)}{(m-1)!} \frac{[L_c - mL_{\text{loop}} - L_p(m)]^m}{m!} e^{-L_c f_m},$$

$$f_m = f_{\text{tw}} + g_{\text{tail}} + l_p(m) \Delta g + m \mathcal{E}_{\text{loop}}, \quad (41)$$

with Λ a cutoff scale for which we choose the helical repeat, as explained in Appendix C where the above expression is derived.

To streamline the notation we define the following densities: the relative linking density

$$r_v := \frac{\nu - \text{lk}}{\rho_{\text{pl}}\omega - \epsilon \text{lk}}, \quad (42a)$$

the relative writhe density

$$r_\omega := \frac{\omega_{\text{loop}}}{\rho_{\text{pl}}\omega - \epsilon \text{lk}}, \quad (42b)$$

the loop writhe density

$$\omega_{\text{loop}} := \frac{W r_{\text{loop}}}{L_{\text{loop}}}, \quad (42c)$$

and the loop density

$$\mu := \frac{m L_{\text{loop}}}{L_c}. \quad (42d)$$

The m -dependent plectoneme length follows as before from the total linking number:

$$l_p(m) = \frac{\nu - \text{lk} - m W r_{\text{loop}} / L_c}{\rho_{\text{pl}}\omega - \epsilon \text{lk}} = r_v - r_\omega \mu. \quad (43)$$

We cannot drop the loop contribution here since we should leave the possibility open that the number of plectonemes increases at the same (or higher) rate as the contour length, reaching some finite density. For the same reason we also keep the end-loop energy. In principle also plectonemes with a negative writhe should be included, but their contribution is very small and in practice only present when tension and linking number are low. We are mainly interested in linking numbers around and above the bifurcation point; thus we can neglect them.

The maximum number of plectonemes can never be higher than L_c/L_{loop} and it is to be expected that finite-size effects easily dominate the turn-extension curves for shorter chains. We want to describe the generic behavior of the turn-extension plot without end effects. The reason is not only to avoid plectoneme-plectoneme interactions, but also to avoid interactions of the magnetic or optical bead with the substrate and details of the exact geometry of attachment of the chain

ends. We write the free energy of the chain as

$$\mathcal{F} = L_c f_0 + m \Delta = L_c \left(f_0 + \frac{\mu}{L_{\text{loop}}} \Delta \right), \quad (44)$$

with Δ as in Eq. (20) and f_0 collecting the terms of the free energy density that do not depend on the loop density.

Assume we are far enough in the plectoneme region that only terms with $m > 1$ contribute. The loop density dependence of the total partition sum reduces to

$$Z \sim \int_0^{\mu_m} d\mu \exp \left\{ \frac{L_c \mu}{L_{\text{loop}}} \left[\ln \left(\frac{l_p(\mu) z(\mu)}{\mu^2} \right) + 2 \ln(L_{\text{loop}}/\Lambda) + 2 - \Delta \right] \right\}, \quad (45)$$

with μ_m the maximum density set by $\mu_m = \sup\{\mu \in [0, 1] | 0 \leq l_p(\mu) \leq 1 - \mu\}$. It is straightforward to verify that the argument of the exponent is a concave function of μ for $\mu \in (0, \mu_m)$ and so its dominant contribution comes from its maximum:

$$\ln \left(\frac{l_p(r_v, \mu) \bar{z}(r_v, \mu)}{\mu^2} \right) - \mu \left(\frac{r_\omega}{l_p(r_v, \mu)} + \frac{1 - r_\omega}{\bar{z}(r_v, \mu)} \right) - \Delta' = 0, \\ \Delta' := \Delta - 2 \ln \left(\frac{L_{\text{loop}}}{\Lambda} \right). \quad (46)$$

Since the relative extension of the chain is $\bar{z} = 1 - r_v - (1 - r_\omega)\mu$ it follows that in the case $r_\omega = 1$ the turn-extension slope does not depend on the number density of plectonemes. Based on our model the value of r_ω is often close to 1 [Fig. 3(d)]. This is one reason why the appearance of multiple plectonemes took so long to discover. The energy per writhe can at the same time differ considerably between loop and plectoneme [$\Delta \neq 0$; Fig. 3(c)], changing the torque after the transition even when the slope can be fitted with just one plectoneme. The more detailed analysis of Eq. (46) is left for Appendix D. Some examples of the dependence of l_p , μ , and \bar{z} as functions of r_v for several combinations of Δ' and r_ω are shown in Fig. 4. The values of Δ' and r_ω corresponding to typical experimental conditions can be read off from Figs. 3(c) and 3(d). It is clear that lowering the salt concentration drives the two strands further apart, thereby decreasing the energetic cost efficiency for writhe production of the plectoneme, and even becoming more costly than the loop itself for low-salt conditions. The formation of plectonemes in that case can be seen as a purely entropic effect. The influence of the tension is slightly more subtle. The tension increases the loop energy $\sim \sqrt{f}$, while in the plectoneme the behavior depends on the salt concentration. At high salt it is only the potential (force) term that changes, since there is not much room for changing the radius, while at

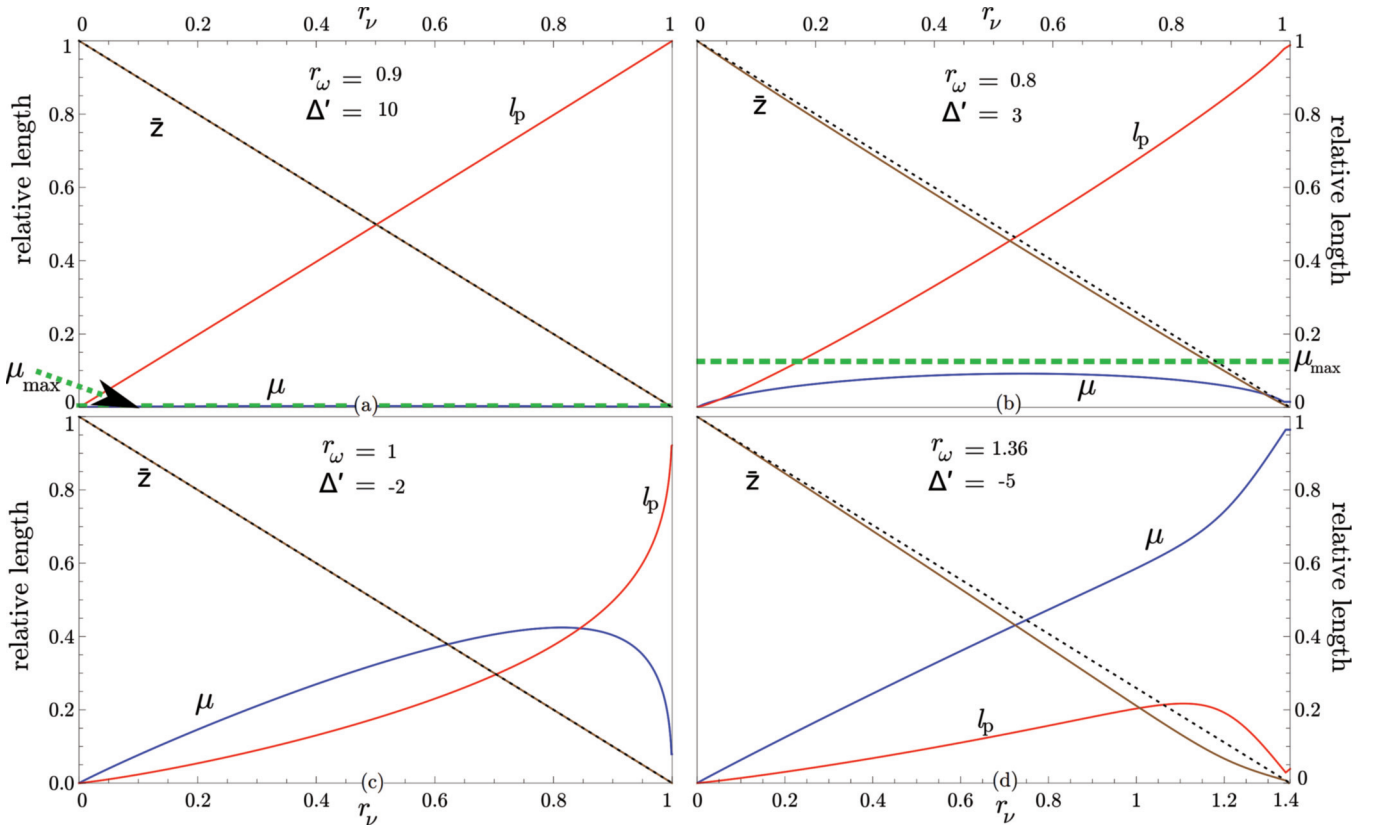


FIG. 4. (Color online) The decrease of the extension \bar{z} and the contributions into which it decomposes (μ and l_p) as functions of scaled linking number density, omitting the straight solution that disappears early on. The plots were generated using the parametrization outlined in Appendix D. The green dashed line is the approximation from Eq. (D7). The dotted line corresponds to the fictive one-plectoneme behavior. (a) Typical single-plectoneme behavior at high cost per writhe difference between loop and plectoneme. (b) Lower Δ increases the number of plectonemes, changing the slope of the turn-extension plot. (c) When the ratio of the writhe densities is 1 the slope does not change even with a large number of plectonemes. (d) When r_ω rises above 1 we end up with a high density of zero-length plectonemes. The force-extension curve also here resembles a one-plectoneme curve but one with modified plectoneme parameters.

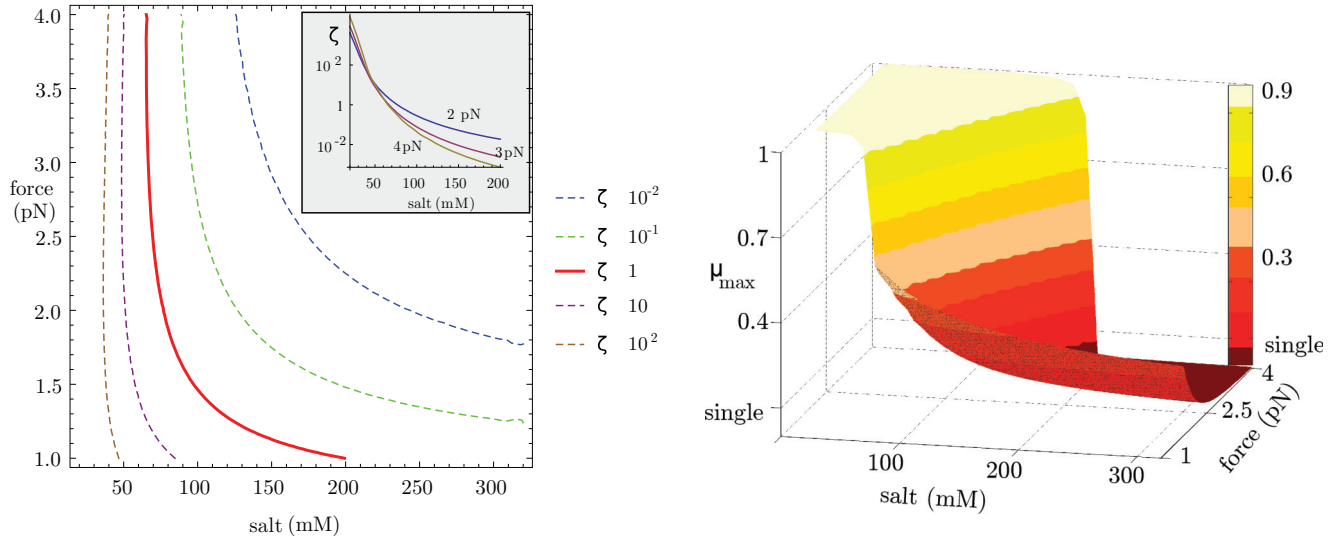


FIG. 5. (Color online) The two faces of multiple plectonemes. (a) Contour plot of the multiplectoneme factor as it depends on salt concentration and tension. The thick red line can be interpreted as the border between a single-plectoneme phase on the right and a multiplectoneme phase on the left. The inset shows ζ as a function of salt concentration for three different forces. Note the crossing of the lines at low salt concentrations. (b) The maximal loop density μ . Note the sharp transition from its maximum ($\mu = 1$) to a vanishing number of plectonemes. Since the maximal μ is reached at the end of the plectoneme slope for $r_\omega > 1$, it does not reflect the multiplectoneme transition along the beginning of the slope.

low salt R has more possibilities to adapt thereby increasing the electrostatic and bending contributions with increasing tension. This is only partly compensated for by an increasing writhe density in the plectoneme. One result of practical use is the multiplectoneme factor ζ :

$$\zeta := r_\omega^2 e^{-\Delta'}. \quad (47)$$

As shown in Appendix D it functions as an indicator for the growth of multiple plectonemes soon after the transition. The value of $\zeta = 1$ it can be interpreted as the boundary between single-plectoneme and multiplectoneme behavior. Its salt and tension dependence is depicted in Fig. 5(a). The largest factor is at low salt and high tension, while for high salt concentrations, ζ increases with decreasing tension. A simpler quantity is the maximal number density for a given tension and salt concentration. Its behavior is depicted in Fig. 5(b). Its change from single-plectoneme to multiplectoneme behavior is also very sharp, but part of this multiplectoneme behavior happens only at the end of the turn-extension plot, when r_ω is larger than 1.

Towards the end of the slope the number density μ goes to zero for r_ω smaller than or equal to 1 while for $r_\omega > 1$ the plectoneme length goes to zero due to the increasing number of plectonemes. This is of course a result of the disappearing of any entropic gain when all of the chain participates in supercoiling. It is interesting to observe that the slopes can for almost all measurements be (often falsely) interpreted as a single-plectoneme slope. When the writhe ratio is above 1, the plectoneme parameters have to be changed, for example by modifying the electrostatic repulsion.

C. Dynamics

At the transition there are two states with equal energy that differ in extension and are separated by an energy barrier. The inclusion of an explicit and realistic loop model allows for an estimate of the transition time from one configuration to the other. Since as argued before the minimal energy path from the straight chain to the plectoneme runs over the family of homoclinic solutions with their free energies given by Eq. (10), we can use the one-dimensional Kramers equation [34], with some adjustment for the nonanalytic potential around the straight configuration, to calculate the transition time between the two states. Although the diffusion coefficients needed to calculate the attempt frequencies are not *a priori* clear, the force dependence can be inferred. The transition times for DNA are at the moment too fast to extract them from available measurements, but with further measurements on the way we plan to come back to this issue in the near future.

We have seen how the appearance of multiple plectonemes can influence the turn-extension curve, but it is often masked by a value of r_ω close to 1. Luckily there is another approach through the torque to which we will come back in the next section. But even the torque behavior after the transition does not necessarily change with the onset of multiple plectonemes. There is yet another property that does always change at the moment that the plectoneme density increases. This is caused by the aforementioned fast twist diffusion: two plectonemes can exchange length through twist-mediated diffusion, which is expected to be much faster than any single plectoneme can diffuse. It also allows for plectoneme diffusion in a crowded environment. This last aspect could be important *in vivo* where for example a change of tension could regulate the “capture” or release of a plectoneme in a pocket within a crowded environment. With this in mind it is interesting to

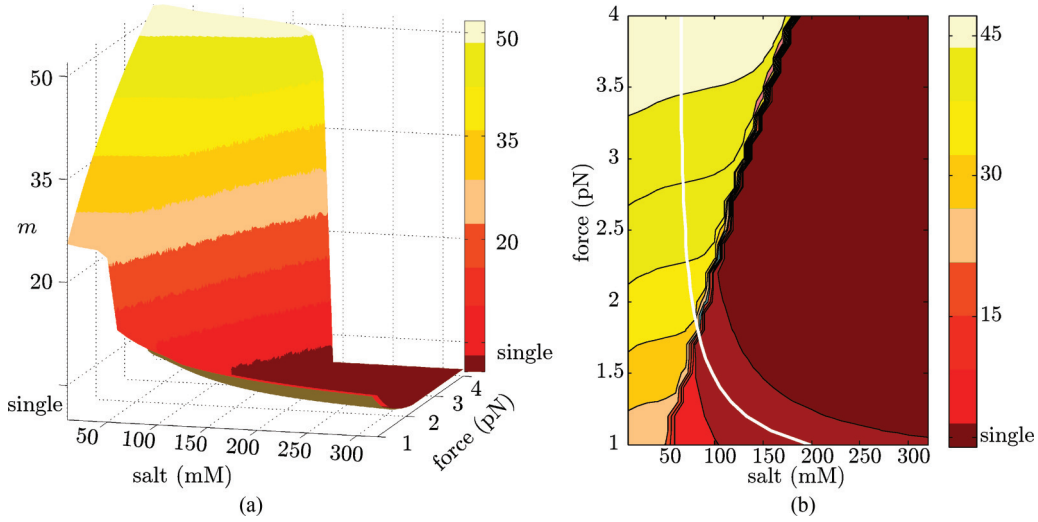


FIG. 6. (Color online) Maximal number of plectonemes as a function of tension and salt concentration for a chain length of 1400 nm. The three-dimensional plot (a) shows how the number of plectonemes goes down with decreasing force at low salt but increases at high salt. The anomalous behavior at low salt is a consequence of the growing loop size limiting the maximal number of loops that fit on the chain. The wiggles are an artifact of the interpolation used. The contour plot of the same data (b) shows a clean border between low and high number. The white line is the $\zeta = 1$ line that marks the border of single- and multiplectoneme behavior. The difference between the two is caused by the growing number of plectonemes at the end of the plectoneme slope, while ζ is a measure for the main part of the slope.

examine the change in the number of plectonemes for a finite chain (Fig. 6). The transition from a single-plectoneme to a multiplectoneme state happens over a narrow band in the tension-salt configuration plane. It again makes sense to speak of two separate phases, the normal single-plectoneme phase characterized by slowly diffusing if not immobile plectonemes and a multiplectoneme phase where plectonemes can diffuse even in crowded environments. It has to be kept in mind that this maximal number of plectonemes might occur only at the end of the plectoneme slope. This is especially true for conditions where $r_\omega > 1$. For this reason ζ might give a better handle on the mobility of plectonemes. It is interesting to note that $\Delta = 0$ plays a key role in the properties of the supercoiling chain as it did in the zero-temperature chain; cf. Eq. (19). The effect of the writhe ratio r_ω , a minor one on ζ but a major one on the maximal loop density, is of entropic origin.

IV. COMPARISON WITH EXPERIMENTS

To test the validity of the model over an extensive range of parameters, use has been made of a series of measurements performed by Seidel and co-workers [22,35]. For combinations of forces from 0.25 to 4 pN and salt concentrations from 20 to 320 mM the turn-extension curves were measured for chains of approximately 600 nm contour length. We smoothed the experimental data with a moving average algorithm. To correct for the geometry of connection to the beads and substrate, the effective contour length of the chain has been obtained by fitting the zero-turn extension to the ideal not-torsional-restricted wormlike chain. Up to lowest order this should be equivalent to the torsionally constrained zero-turn configuration. The effective chains thus obtained have a length that varies between 570 and 630 nm. A set of measurements under varying forces but constant salt concentration has been performed on one chain, allowing us to verify that the effective

chain length stays more or less constant once the geometry of the chain attachment is fixed. Only for forces below 1 pN does the effective chain length decrease. This is partly due to the bent chain attachment, combined with chains too wildly fluctuating for our perturbative model.

The minimization procedure was initiated as follows: starting from the bifurcation point lk_{cr} , the applied linking number per length was set to $\nu = lk_{cr} + 0.2$ to assure that the linking number density is far after the transition. The parameters of the model were set to $lk = 0.8lk_{cr}$, $\alpha = 1$, $\sigma_r = 1/(2\kappa)$, and $R = 1 + \kappa^{-1}$ (in nanometers such that the potential of a cylinder with a radius of 1 nm is in the linear regime at R). The free energy for a single chain was minimized after which the obtained values were used to set ν to $(\omega + lk_{str} + 2lk_{cr})/2$, setting the linking number density halfway between the bifurcation point and the maximum. The resulting plectoneme parameters were used as starting values for another minimization. In that way the applied linking number is approximately halfway between the critical value and the maximal value. The reasoning is that with a linking number close to the bifurcation point the influence of an incomplete description of the end loop becomes too strong, while a linking number too far from the transition might underestimate the influence of multiplectoneme configurations. The precise value is not very important. Too close to the bifurcation point the chain collapses before the transition in low-salt conditions. The reason is not so much the influence of the loop but a K^2 value that gets too low. Of course any prediction based on the model for K^2 values below 3 is unreliable. The generation of the force-extension behavior is based on plectoneme energies from this minimization. The whole procedure is very fast.

As a first test of our model we compare predicted plectoneme slopes to those determined in experiments. Note that the choice of where to measure the slope is not always obvious in either theory or experiment. Whenever there was a clear

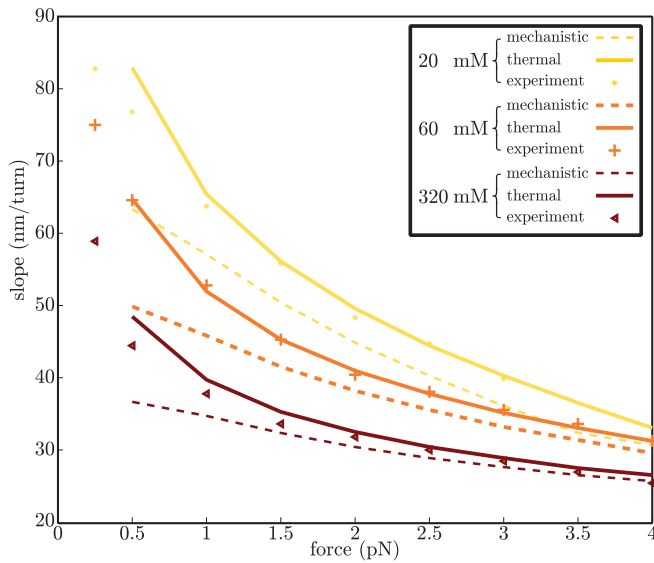


FIG. 7. (Color online) Slopes with and without thermal contributions. The dotted curves were calculated from the model as described up to Sec. III, Eq. (17). The solid thermal curves include multiplectonemes and were calculated using the method outlined in the text.

constant slope visible it was taken as the slope; otherwise the first slope after the transition was taken. Especially for the short 600 nm chains it was not always clear what to take as slope. This is especially true for low salt, 20 to 60 mM, conditions. Nonetheless the slopes for the full range indicated a nice agreement between experiment and model. The results for 20, 60, and 320 mM are in Fig. 7. The influence of the multiplectoneme phase is clearly visible for low salt concentrations. There is also a clear improvement in the low-force range, although there the value of K^2 of 2 or lower around the transition point makes the agreement merely coincidental. The turn-extension plot at 20 mM and 3 pN in Fig. 8(a) shows

the details. The transition happens at a lower linking number than in the experiment, presumably because it is too close to the bifurcation point for a reliable perturbative calculation. To produce the plots the torsional persistence length was lowered to 90 nm from 110 nm to get the transition point close to the experimental value. In Fig. 8(b) the number of plectonemes is set out against the number of turns for these conditions.

The curves for 20 and 320 mM are shown in Fig. 9. For most cases our model predicts the experimental curves well. The behavior at the transition at 20 mM and a tension above 3 pN is not well defined perturbatively, since the straight solution has a K^2 value below 3 before the first plectoneme solution becomes available. The 20 mM measurements show an exceptional behavior at 3.5 pN. It is possible that the chain undergoes a phase transition as has been suggested [36]. Another possibility is that because plectoneme formation is relatively expensive, starting plectonemes are extremely unstable. That can explain the sawtooth behavior with signs of attempts at plectoneme nucleation.

A set of experiments performed on a 3850 nm chain in a 320 mM solution with the same setup shows a longer clear slope in Fig. 10(a). The transition point suggests here a 120 nm torsional persistence length.

Another test of the model is the analysis of the plectoneme torques. The torque is obtained by dividing the increase of the free energy by the rotation angle that caused it. It is commonly believed that the linear slope of the curves coincides with a state of constant torque [38,39]. This makes it attractive to use the DNA plectoneme as a source of constant torque in the study of molecules that interact with DNA, like topoisomerase and helicase. One way to measure this plectoneme torque is by using a specially nanofabricated quartz cylinder in conjunction with an optical tweezer [40]. The setup seems to be very promising, enabling the measurement of torque at the same time as force and extension. A small set of measurements were done with relatively short chains of 700 nm [41]. Another method makes use of the constant torque in the plectoneme

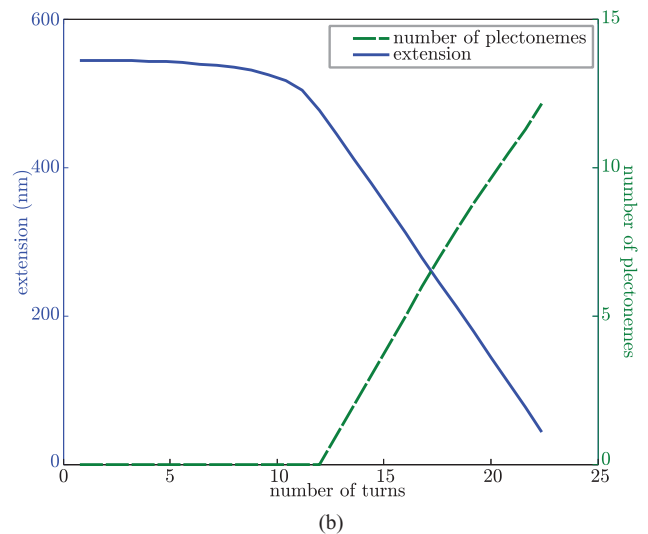
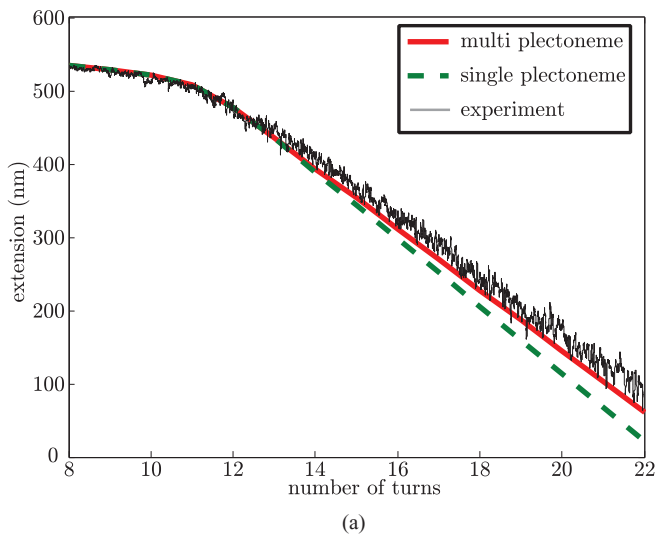


FIG. 8. (Color online) Influence of multiplectonemes on the turn-extension plot at low salt. For a chain with a contour length of 600 nm and a tension of 3 pN at a 20 mM salt concentration there is a noticeable effect on the slope (a). The growing number of plectonemes and not a growing plectoneme length is responsible for the slope in (b). The experimental data are from Seidel and co-workers [22,35].

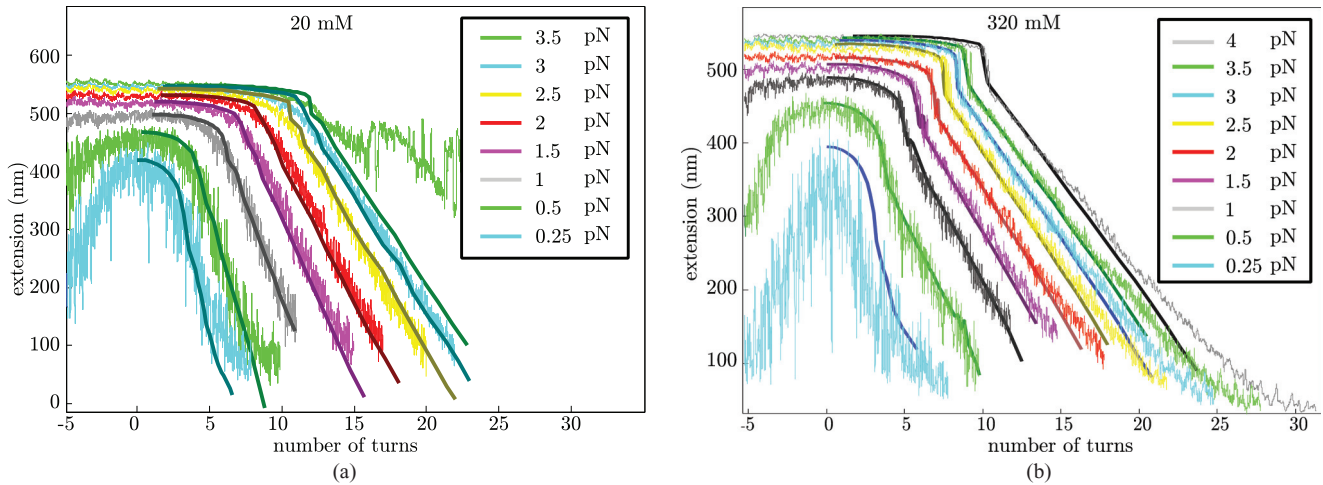


FIG. 9. (Color online) Turns versus extension plots for 20 mM (a) and 320 mM (b) monovalent salt concentrations under varying tension. The torsional persistence length for most salt gave the best fit for 110 nm. For 20 mM a lower value of 90 nm had to be taken to get an almost perfect agreement, but as explained in the text it might be a calculational artifact due to the proximity of the bifurcation point.

region combined with Maxwell relations between torque and linking number and force and extension as free energy parameters. The method calculates the plectoneme torque over a large range of forces using an approximately linear linking number–torque relation before the transition at high tensions. Assuming a constant torque after plectoneme formation, the torque for a large range of data can be calculated just from the turn–extension plot. This is the setup from Mosconi *et al.* [42]. The resulting torques in the two types of measurements [41,42] seem to differ. It could be that the salt concentrations differ too much, or that the response of the optical trap is too slow. It is interesting to compare the torques that our model predicts with those of Ref. [42]. To our surprise the torques we calculate differ from their measurements substantially enough to cast doubt on the validity of our model; see Table I. The torque from the model was calculated just after the transition at the start of the plateau by calculating the change in free energy as a function of the change in linking number. This deviation in torque was not totally unexpected, since the torque data were one of the reasons for Maffeo *et al.* [22] to incorporate a charge reduction factor into their model. What is somewhat mysterious is that the force–extension curves themselves are in good agreement with our model as illustrated by Fig. 10(b).

TABLE I. Indirect torque measurements using Maxwell relations [42] compared to the theoretical values from our model

Salt (mM)	Force (pN)	Expt. torque (pN nm) [42]	Theoretical torque (pN nm)
10	2.86	28.1	35.0
	2.53	26.2	32.0
50	3.66	29.6	34.7
	3.23	27.4	32.4
100	3.33	24.4	30.1
	2.61	20.7	26.3
500	4.33	22.3	29.6
	3.80	20.2	27.5

If the torque depends only on the shape of that curve, while Maxwell’s relations hold per definition, somewhere a wrong assumption must have been made.

Comparing our torque predictions with the direct torque measurements from the older optical tweezer measurements [41] reveals however a remarkably good agreement as is shown in Fig. 11(b), where the torques are shown as a function of the supercoiling density defined as the ratio of the linking number density to the linking number density of the two strands of the double helix when the chain is straight and relaxed. This last density is of course $1/\text{helical repeat} = 1/3.6 \text{ nm}^{-1}$.

The culprit is readily revealed as the multiplectoneme phase. In extracting the torque from the turn–extension curves an essential assumption is that the torque in the linear slope is constant. That almost presupposes that the slope is a one-plectoneme slope. Lacking a method to verify this assumption, it had to be accepted on face value. In reality the torque is not constant at all for lower forces. Thanks to the rapidly increasing number of plectonemes along the chain, the torque is almost linearly *increasing*, invalidating the calculations. When we take this increase into account, the resulting torque values agree again wonderfully well with the predictions from our model.

As an example we borrow the calculations from Mosconi *et al.* [37]. The relevant curves are in Fig. 12. The Maxwell relation calculations are performed over the path as shown in the figure on the left. The resulting torque for 3.67 pN is 27 pN nm. But if we examine the torque as calculated from the model the result is higher, around 34.9 pN nm. Although the torque is constant for the high-tension slope, the path from B to C in Fig. 12 is one of decreasing torque, thereby resulting in a too low estimate for the plectoneme torque.

Inspired by our preliminary results an experiment was set up for which the first results [43] became recently available. Multiplectonemes were visualized under conditions where we had expected them to exist. Also the fast twist-mediated diffusion, only possible when at least two plectonemes are present, was observed. The resolution is at this moment not good enough to extract plectoneme number densities.

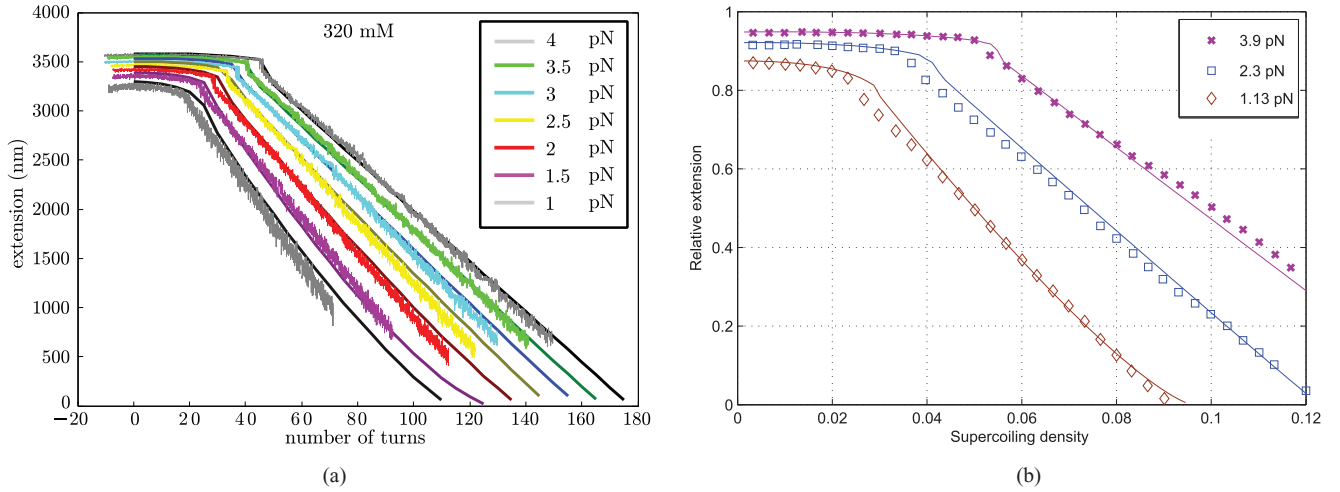


FIG. 10. (Color online) Long chains. (a) Turns versus extension plots for a chain of 3540 nm with tension from 1 to 4 pN in 320 mM salt. A torsional persistence length of 120 nm was used. (b) Some of the magnetic tweezer measurements from Mosconi *et al.* [37] with a 5.4- μm -long chain in 100 mM salt for three different forces. The curves are from our model with a torsional persistence length of 115 nm.

V. COMPARISON WITH OTHER MODELS AND OUTLOOK

Over the years numerous models have been proposed, all of them bringing in some new ideas. We cannot compare our model against all of them, but we will discuss some recent works that are of interest with respect to our model.

First of all there is the model proposed by Seidel and co-workers [22] where, based on a nonthermal model like the one in Sec. II, it was proposed that the charge of the DNA molecule should be reduced by a factor with a value determined by the experimental slopes. The reasoning was that some of the counterions might be confined to the grooves of the double helix. An assumption was that the thermal shortening of the DNA would be the same in the plectoneme and in the tails. Finally the resulting potential was used as the basis of Monte Carlo (MC) simulations that confirmed

that the slopes were unaffected by fluctuations. There exist a couple of objections. The lack of any influence of thermal fluctuations is hard to understand, since the way fluctuations are restricted is quite different in tails and plectoneme. It might be that the 5 nm segment length chosen for the MC simulations is too large for capturing the essential part of the spectrum. Perhaps more problematic is the charge reduction of more than one-half. It would mean that the concept that DNA is a strong polyelectrolyte with respect to the phosphates, one charge per base pair, is wrong. This would contradict direct experimental evidence (for example Ref. [44]), but also indirect measurements, e.g., concerning the pressure of viral DNA, DNA condensation models, and more. It is puzzling how the inhomogeneity of small monovalent counterions far within the inner layer can affect the potential outside the nonlinear domain. Furthermore as we have shown the slopes are in fact

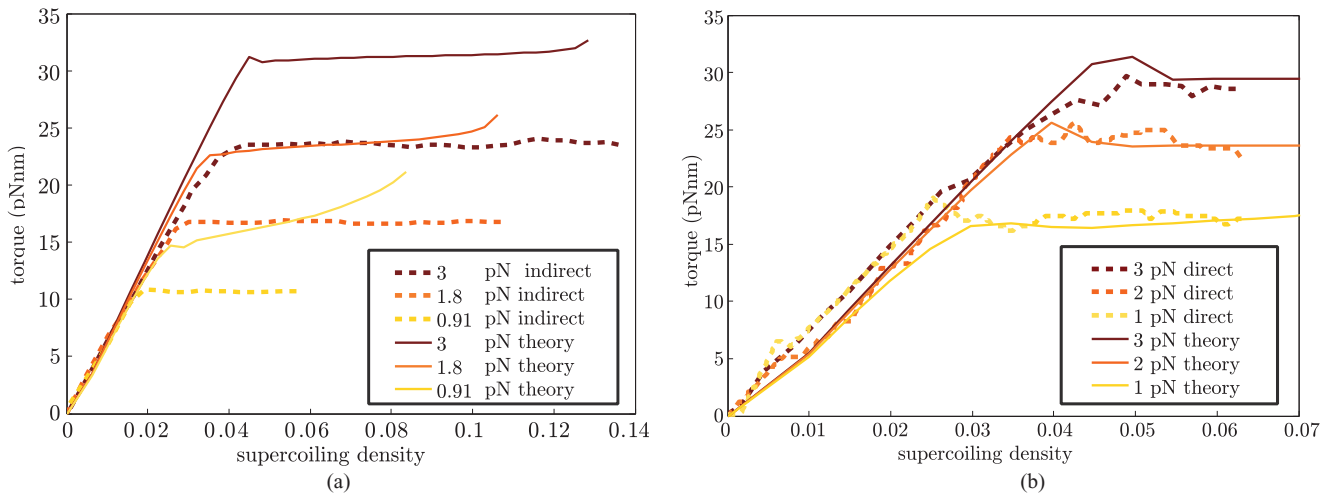


FIG. 11. (Color online) (a) Torque as a function of supercoiling density for three different tensions calculated from the model compared to the indirect determination of the torque using Maxwell relations under the assumption of constant plectoneme torque [37]. The conditions are the same as in Fig. 12. (b) Comparison between experimental results from an optical tweezer experiment using a quartz cylinder to directly measure the torque [41] and our theoretical model. The DNA molecule has a contour length of 725 nm. The monovalent salt concentration is 150 mM.

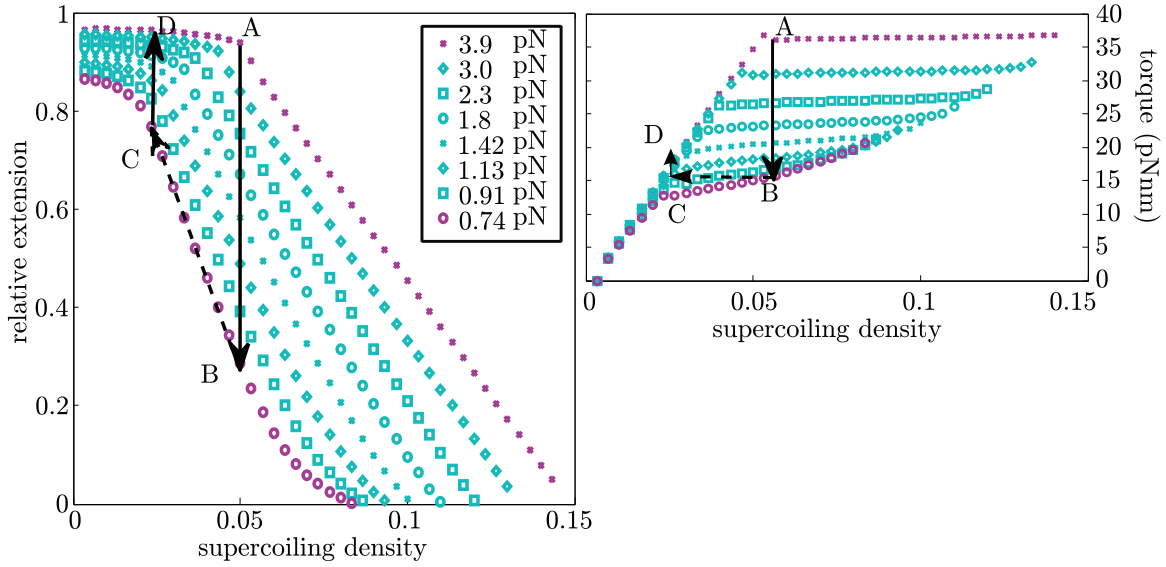


FIG. 12. (Color online) Illustration of the magnetic tweezer measurements from Mosconi *et al.* [37] as a basis for indirect torque measurements. The curves are calculated from our model for a range of tensions where fluctuations are small enough, using the criterion $K^2 \geq 3$. The contour length is $5.6 \mu\text{m}$ and the monovalent salt concentration is 100 mM. On the left are the resulting turn-extension plots. On the right are torques from our model. Notice how the torque is increasing in the plectoneme region, caused by the growing number of plectonemes. The circuit *ABCD* results in too high a torque at *C* when one assume the plectoneme torque to be constant.

not very dramatically affected since the writhe ratio is close to 1. Another more recent work introduced small loops in addition to the possibility of multiple-plectoneme formation [33]. The resulting modeling cannot faithfully reproduce the measured curves though. One problem is that these little loops do not have any electrostatic or entropic repulsion incorporated. It is possible to add a more detailed description to these loops, as we had considered in an early version of this work that was presented at the CECAM workshop “Coarse grain mechanics of DNA: bases to chromosomes” in 2010 which took place in Lyon [45], but then it is only a small step to acknowledge that these loops are in fact zero-length plectonemes.

In this paper we have developed a model for plectoneme formation that makes a consistent description of thermal fluctuations on all scales. The model is perturbative, which has its limitations, but under conditions where a perturbative treatment makes sense it performs well. We discovered a sharp boundary between single-plectoneme and multiplectoneme behavior and argued for possible biological implications. This multiplectoneme behavior at the same time resolves some anomalies in experimental data. There are of course still many open questions. We think that the most pressing concerns the direct measurement of torques over the full plectoneme slope. This could shed some light on the correct cutoff value. A full analysis of the length drop at the transition we will leave for a future presentation.

ACKNOWLEDGMENTS

This research would have been impossible without the excellent sets of measurements provided by the group of Ralf Seidel. We are thankful for the inspirational discussions with Theo Odijk and Andrey Cherstvy. This work would not have been possible without the support from Gerhard Gompper

and the Forschungszentrum Jülich. We acknowledge the many discussions with Marijn van Loenhout and Cees Dekker and are glad that our theoretical predictions helped them to find the multiplectoneme phase experimentally.

APPENDIX A: WRITHE OF A PLECTONEME

In principle the writhe of the plectoneme can be calculated using Fuller’s equation and continuity. Care should be taken since the plectoneme moves through a curve with an antialigned tangent once every full turn of the plectoneme, when one considers the (un)winding as the homotopy to the straight line. Since we intend to use an exact expression for the writhe at least for the ground state it is instructive first to calculate the writhe density for the plectoneme using Fuller’s equation with respect to the plectoneme axis for both strands, forgetting loop and tail:

$$\begin{aligned}\omega_1(s) &= \frac{1}{2\pi} \frac{\cos(\alpha)(\sin(\alpha) - 1)}{R(t)}, \quad s \in [0, l_p/2], \\ \omega_2(s) &= \frac{1}{2\pi} \frac{\cos(\alpha)(\sin(\alpha) + 1)}{R(t)}, \quad s \in [l_p/2 + l_l, l_p + l_l].\end{aligned}\quad (\text{A1})$$

We could in a hand-waving fashion define an “average” writhe density as

$$\omega(\alpha, t) \stackrel{?}{=} \frac{1}{2} [\omega_1(s) + \omega_2(l_p + l_l - s)] = \frac{\cos(\alpha) \sin(\alpha)}{2\pi R(t)}. \quad (\text{A2})$$

The problem is that this definition, giving the usual relation, is based on Fuller’s equation with respect to another axis than the one we started with, and we have not taken the writhe of the end loop into account.

A correct way that shows the importance of the rotation of the closing loop is to use the z axis as reference here also. Opposing points on the plectoneme strands have in this case

the same writhe:

$$\omega_b(s) = \omega_b(l_p + l_l - s) = \frac{1}{2\pi} \frac{\sin \alpha \cos \alpha}{R(t)} \times \left[1 - \frac{1}{1 + \cos \alpha \cos \left((s_0 + s) \frac{\cos \alpha}{R(t)} \right)} \right]. \quad (\text{A3})$$

A surprising s dependence enters the writhe density of the plectoneme. The subscript b is as a reminder that this is a bare writhe density that does not include interactions with the rest of the chain. Adding plectoneme length also changes the writhe of the end loop though. The closing end loop is described at the onset of the plectoneme formation by some space curve $\mathbf{r}_0(u) = (r_x(u), r_y(u), r_z(u))$, $u \in [0, l_l]$, with boundary conditions $\mathbf{r}_0(0) = \mathbf{r}_p(0)$ and $\mathbf{r}_0(l_l) = \mathbf{r}_p(l_l)$. We furthermore assume the connection between the plectoneme and the end loop to be smooth, making the tangent well defined at the boundaries. The increase of the plectoneme by an amount of contour length $2s$ causes the end loop to rotate around the x axis by an angle $\phi(s) = s \cos \alpha / R(t)$. The rotated loop is given by

$$\mathbf{r}_s(u) = \hat{O}_x(\phi(s))\mathbf{r}_0(u) = \begin{pmatrix} r_x(u) \\ \cos \phi(s)r_y(u) + \sin \phi(s)r_z(u) \\ -\sin \phi(s)r_y(u) + \cos \phi(s)r_z(u) \end{pmatrix}. \quad (\text{A4})$$

This rotation induces an s -dependent change in the writhe of the loop to

$$\begin{aligned} \text{Wr}_{\text{loop}}^1(s) &= \frac{1}{2\pi} \int_0^{l_l} du \left(\frac{\cos \phi(s)[t_x(u)\dot{t}_y(u) - \dot{t}_x(u)t_y(u)]}{1 - \sin \phi(s)t_y(u) + \cos \phi(s)t_z(u)} \right. \\ &\quad \left. - \frac{\sin \phi(s)[t_z(u)\dot{t}_x(u) - \dot{t}_z(u)t_x(u)]}{1 - \sin \phi(s)t_y(u) + \cos \phi(s)t_z(u)} \right). \end{aligned} \quad (\text{A5})$$

The superscript is just a reminder that it is not the writhe of the full homoclinic solution, but just of that part that detaches to function as end loop for the plectoneme. Note that this writhe is not necessarily well defined. In fact since the length of the loop is finite, its x component is bounded and thus has at least one point where the tangent lies in a plane perpendicular to the x axis. This tangent will be once every full turn of the plectoneme antipodal to the z axis and thus invalidates Fuller's equation.

We can nonetheless calculate the differential change of this writhe per plectoneme contour:

$$\begin{aligned} \frac{d\text{Wr}_{\text{loop}}^1}{ds} &= \frac{\cos \alpha}{2\pi R(t)} \int_0^{l_l} du \\ &\quad \times \frac{-\dot{t}_x(u) - \sin \left(s \frac{\cos \alpha}{R(t)} \right) [t_x(u)\dot{t}_y(u) - \dot{t}_x(u)t_y(u)]}{\left[1 - \sin \left(s \frac{\cos \alpha}{R(t)} \right) t_y(u) + \cos \left(s \frac{\cos \alpha}{R(t)} \right) t_z(u) \right]^2} \\ &\quad - \frac{\cos \left(s \frac{\cos \alpha}{R(t)} \right) [t_z(u)\dot{t}_x(u) - \dot{t}_z(u)t_x(u)]}{\left[1 - \sin \left(s \frac{\cos \alpha}{R(t)} \right) t_y(u) + \cos \left(s \frac{\cos \alpha}{R(t)} \right) t_z(u) \right]^2} \\ &= \frac{\cos \alpha}{\pi R(t)} \frac{t_x(0)}{1 - \sin \left(s \frac{\cos \alpha}{R(t)} \right) t_y(0) + \cos \left(s \frac{\cos \alpha}{R(t)} \right) t_z(0)}, \end{aligned} \quad (\text{A6})$$

where use has been made of the unimodularity of the tangent vector and its symmetry: $t_x(0) = -t_x(l_l)$, $t_{y,z}(0) = t_{y,z}(l_l)$. Making use of the boundary conditions we finally find

$$\frac{d\text{Wr}_{\text{loop}}^1}{ds} = \frac{\cos \alpha \sin \alpha}{\pi R(t)} \frac{1}{1 + \cos \alpha \cos \left((s_0 + s) \frac{\cos \alpha}{R(t)} \right)}. \quad (\text{A7})$$

By adding this differential writhe density to the “bare” writhe density of the plectoneme as given by Eq. (A3) (half of it to each strand) we recover the standard writhe density of a plectoneme (16), but now with the added bonus that the remaining writhe of the end loop is independent of the length of the plectoneme. Since it is only in the end loop that antipodal points appear along the homotopy, defined by the explicit formation of the plectoneme, we can state that in this sense the writhe is additive:

$$\text{Wr}(t, \alpha) = \text{Wr}_{\text{loop}}(t) + L_p \omega(t, \alpha), \quad (\text{A8})$$

with Wr_{loop} and ω given by Eqs. (9) and (16).

Note that we implicitly used continuity to recover the full writhe of the chain by adding the differential writhe change of the end loop. In hindsight it is clear that the end loop should be included in the final result. Imagine, for example, a larger end loop such that the helices do not intertwine. The writhe in this case can be calculated immediately without any continuity argument and it is easy to show from Eq. (A3) that applying Fuller's equation to a chain with such a nonintertwining plectoneme of n turns [$l_p = 4n\pi R(t)/\cos \alpha$] gives a writhe of $\text{Wr} - 2n$.

APPENDIX B: FLUCTUATIONS OF THE STRANDS IN A PLECTONEME

Our treatment of thermal fluctuations in the plectoneme follows largely the work by Ubbink and Odijk [17], with some change forced upon us by the physical conditions. In our case we cannot just use Burkhardt's result for the confinement of a rotational relaxed chain [46], but have to take the twist along the chain into account. This has two implications: (1) The confinement free energy gets twist-dependent corrections, the calculation of which are presented in Ref. [30]. (2) The confinement gives a relation between linking number and twist that depends on the confinement channel width. This is used to calculate the contour length of the plectoneme in the text.

In this Appendix we will discuss how the fluctuations can be separated from the average plectoneme path. Thermal undulations effectively shorten the chain within its superhelical path. This has implications for the bending energy and the writhe density of the plectoneme. To calculate the effect we attach to each point along the nonundulating path, the zero path or zero chain, a triad, consisting of the tangent at that point and two normals. The fluctuations we can express as deviations in the two normal directions from the zero path. The deviations in the tangential direction follow from the inextensibility, or if needed a finite stretch modulus can be included [26]. For the plectoneme as triad we take its Fresnet basis, where the normal is the direction of curvature, which is the radial direction, making the “pitch direction” the binormal. With respect to the

contour length the point along the zero path gets shifted by a shortening factor ρ , for which we will use its expectation value. The deflection length in a confined channel is considerably shorter than the persistence length of the chain. In general one can expect, in conditions that allow for a perturbative expansion, that the length scales of the fluctuations are small compared to the global length scales. The main assumption in the following is that the wavelength of thermal undulation is considerably shorter than those of the writhing zero path. More precisely the curvature and Fresnet torsion, which is 2π times the writhe density of the plectoneme, are small compared to the wave numbers of thermal undulations. Neglecting contributions from the zero-path torsion and curvature we arrive at the following equations:

$$\begin{aligned} \mathbf{r}(s) &:= \mathbf{r}_0(\rho s) + u_i(s) \mathbf{t}_{\perp,0}^i(\rho s) \\ \Rightarrow \mathbf{t}(s) &\simeq \rho \mathbf{t}_0(\rho s) + \dot{u}_i(s) \mathbf{t}_{\perp,0}^i(\rho s) \\ \Rightarrow \dot{\mathbf{t}}(s) &\simeq \rho^2 \dot{\mathbf{t}}_0(\rho s) + \ddot{u}_i(s) \mathbf{t}_{\perp,0}^i(\rho s). \end{aligned} \quad (\text{B1})$$

We conclude that we can treat the channel as being straight for thermal fluctuations, provided we multiply the curvature of the zero chain by ρ^2 . The bending energy of the zero chain, being proportional to the curvature squared, acquires then a factor of ρ^4 .

For the writhe calculation we make again use of Fuller's equation (6), now with a homotopy from the zero path. It is fairly easy to prove, using short intervals and continuity, that the projection of the fluctuating path on the zero path along the normal bundle forms a valid homotopy for Fullers equation. We write $\omega(s) = \omega_0(\rho s) + \Delta\omega(s)$. The zero-path writhe is as before but multiplied by ρ , since the tangent at ρs does not change, but its rate of change does.

Finally applying Fuller's equation results in

$$\begin{aligned} \Delta\omega(s) &= \frac{1}{2\pi} \frac{[\mathbf{t}_0(\rho s) \wedge \mathbf{t}(s)][\dot{\mathbf{t}}(s) + \rho \dot{\mathbf{t}}(\rho s)]}{1 + \rho} \\ &\simeq \frac{1}{4\pi} [\dot{u}_r(s) \ddot{u}_p(s) - \ddot{u}_r(s) \dot{u}_p(s)] \end{aligned} \quad (\text{B2})$$

up to quadratic order and using the same assumptions as before.

APPENDIX C: MULTIPLECTONEME ENTROPY

In this Appendix the number of configurations of a chain with a total plectoneme length L_p , divided over m plectonemes, $Z_m(L_c, \text{lk})$, is calculated. We make lengths dimensionless by rescaling them with a cutoff. The natural cutoff is not *a priori* clear. One could argue for the deflection length λ , which is the natural length scale in the tails, or alternatively for the 3.5 nm helical repeat, which must be a scale that influences nucleation of loops. We will choose the latter as the length scale for positioning and length distribution of the plectonemes in our calculations. Measurement data are, due to noise, not yet precise enough to differentiate between possible length scales. Fluctuations in plectoneme length L are mostly balanced by twist fluctuations. Their contribution to the partition sum is independent of the way L is split between plectonemes, and thus

$$Z_m(L_c, \nu) = \int_0^{L_c - mL_{\text{loop}}} dL z_m(L) \exp[-L_c f_m(L, \nu)] \quad (\text{C1})$$

with $z_m(L)$ the density of states at constant L . We assume that $z_m(L) \simeq z_m(L_p)$, constant over the sharply peaked minimum of $f_m(L, \nu)$ around $L = L_p$. From Eqs. (43) and (24) and because of the sharp minimum we find that the integral can be approximated by a Gaussian:

$$Z_m(\text{lk}) = \sqrt{\frac{L_c}{2\pi\rho_{\text{pl}}^2 P_c^{\text{ren}}}} z_m(L_p) \exp[-L_c f_m(L_p, \text{lk})]. \quad (\text{C2})$$

We first treat the case with hard-core interactions between the plectonemes. For a configuration with one plectoneme of length L_p and loop length L_{loop} the number of possible configurations is $L_c - L_{\text{loop}} - L_p$, the length along the chain where the plectoneme can end. In case of two plectonemes sharing the length L_p , the first plectoneme we encounter, with plectoneme length Λ_1 , can have a position x_1 between $L_{\text{loop}} + \Lambda_1$ and $L_c - L_{\text{loop}} - (L_p - \Lambda_2)$, while the second plectoneme can have a position x_2 in the interval $[x_1 + \Lambda_2 + L_{\text{loop}}, L_c]$. It is easy to show using induction that the partition sum for m loops can be written as

$$\begin{aligned} z_m^{\text{hc}}(L_p) &= \prod_{i=1}^{m-1} \left(\int_0^{L_p - \sum_{j=0}^{i-1} \Lambda_j} d\Lambda_i \right) \\ &\times \prod_{k=1}^m \left(\int_{x_{k-1} + L_{\text{loop}} + \Lambda_k}^{L_c - (m-k)L_{\text{loop}} - (L_p - \sum_{q=1}^k \Lambda_q)} dx_k \right), \end{aligned} \quad (\text{C3})$$

with $\Lambda_0 = x_0 = 0$. To shorten the notation we define an effective chain length $L'_c := L_c - mL_{\text{loop}}$. The second product, which we denote by y_m , integrates over all positions of the plectoneme. It can be written as

$$\begin{aligned} y_m(L'_c - L_p) &= \prod_{k=1}^m \left(\int_0^{L'_c - L_p - \sum_{q=0}^{k-1} x_q} dx_k \right) \\ &= \int_0^{L'_c - L_p} dx_1 y_{m-1}(L'_c - L_p - x_1) \\ &= \mathcal{L}^{-1} \left(\frac{1}{t^{m+1}} \right) (L'_c - L_p) \end{aligned} \quad (\text{C4})$$

$$= \frac{(L'_c - L_p)^m}{m!}, \quad (\text{C5})$$

where in the third step \mathcal{L}^{-1} denotes an inverse Laplace transform and the *faltung* theorem has been used. The first term can be calculated analogously, resulting in the comprehensive result

$$Z_m^{\text{hc}}(L_c, \nu) \sim \sqrt{L_c} \frac{L_p^{m-1}}{(m-1)!} \frac{(L'_c - L_p)^m}{m!} e^{-L_c f_m}. \quad (\text{C6})$$

This hard-core interaction is probably not entirely realistic. With a minor penalty plectonemes can have some overlap. The effects of plectoneme interactions come into play only when most of the free DNA has been used. As a test the calculations can be performed with the other extreme of noninteracting plectonemes. Defining $L'_c := L_c - L_{\text{loop}}$, and again implicitly rescaling all lengths by the helical repeat, we

find as combinatorial factor

$$\begin{aligned} z_m^{\text{ni}}(L_p) &= \frac{1}{m!} \prod_{i=1}^m \left(\int_0^{L_p - x_{i-1}} dx_i (L_c'' - x_i) \right) \\ &\quad \times \left(L_c'' - L_p + \sum_{i=1}^{m-1} x_i \right) \\ &= \frac{1}{m!} \mathcal{L}^{-1} \left(\frac{L_c''}{t} - \frac{1}{t^2} \right)^m (L_p) \\ &= \frac{1}{m!} \sum_{k=0}^m \binom{m}{k} \frac{(-1)^k L_c''^{m-k} L_p^{m+k-1}}{(m+k-1)!}, \quad (\text{C7}) \end{aligned}$$

which, not necessarily providing more clarity, can be written using a confluent hypergeometric function as

$$Z_m^{\text{ni}}(L_c, v) \sim \sqrt{L_c} \frac{(L_c'')^m L_p^{m-1} {}_1F_1(-m, m, L_p/L_c'')}{m!(m-1)!} e^{-L_c' f_m}. \quad (\text{C8})$$

The experiments we have analyzed are mostly situated in the relatively low plectoneme density range, making the

difference between these two extremes too small to be able to tell how soft the plectoneme interaction is, especially in light of the fact that the relative writhe densities of plectoneme and loop do not differ much.

APPENDIX D: ANALYZING THE PLECTONEME LENGTH AND NUMBER DENSITY

In this Appendix we analyze the behavior of the plectoneme length and number density as the number of turns increases. Note that Eq. (46) is symmetric under the transformation $r_n, r_\omega \rightarrow 1 - r_n, 1 - r_\omega$ whereby $l_p \leftrightarrow \bar{z}$. The question is when to expect multiple plectonemes. Defining $\mu_l := \mu/l_p$ and $\mu_z := \mu/\bar{z}$, we can formally solve Eq. (46), exponentiating it as

$$r_\omega \mu_l \exp(r_\omega \mu_l) = \exp[-(1 - r_\omega) \mu_z - \Delta'] \frac{r_\omega}{\mu_z}. \quad (\text{D1})$$

This is the defining equation for the Lambert W function [47]. Since the right-hand side is positive we need the principal branch W_p as the only real-valued branch:

$$\mu_l = \frac{1}{r_\omega} W_p \left(\frac{r_\omega}{\mu_z} e^{-(1-r_\omega)\mu_z - \Delta'} \right), \quad (\text{D2a})$$

$$\mu_z = \begin{cases} \frac{1}{1-r_\omega} W_p \left(\frac{(1-r_\omega)}{\mu_l} e^{-r_\omega \mu_l - \Delta'} \right) & \text{if } r_\omega < 1 \text{ or } r_\omega > 1 \wedge \mu_z < \frac{1}{r_\omega - 1}, \\ \frac{1}{1-r_\omega} W_m \left(\frac{(1-r_\omega)}{\mu_l} e^{-r_\omega \mu_l - \Delta'} \right) & \text{if } r_\omega > 1 \wedge \mu_z > \frac{1}{r_\omega - 1}, \\ \frac{1}{\mu_l} e^{-\mu_l - \Delta'} & \text{if } r_\omega = 1. \end{cases} \quad (\text{D2b})$$

The second equation follows from the first by symmetry, but care has to be taken which branch to follow. The argument is in this case negative and a second real branch exists, W_m . The crossover happens when the argument reaches its minimum of $-1/e$.

We next analyze the scaling behavior at the end of the plectoneme slope, when \bar{z} reaches zero, using Eq. (D2b).

Case $r_\omega < 1$. Suppose we end up with a finite density of plectonemes; then $\mu_z \rightarrow \infty$ and so the argument of $W_p \rightarrow \infty$ in Eq. (D2b). As this is impossible, we have $\mu \rightarrow 0$ and $l_p \rightarrow 1$. Since μ_l tends to zero the argument of W_p goes to infinity. To lowest order we find

$$\begin{aligned} \lim_{\bar{z} \rightarrow 0} \mu_z &\simeq \frac{1}{1-r_\omega} \ln \left[\frac{1-r_\omega}{\mu_l} e^{-(1-r_\omega)\mu_l - \Delta'} \right] \\ &\simeq -\frac{1}{1-r_\omega} \ln(\mu) \\ \Rightarrow \lim_{\bar{z} \rightarrow 0} \mu &\simeq \frac{\bar{z}}{1-r_\omega} W_p \left(\frac{1-r_\omega}{\bar{z}} \right) \simeq -\frac{\bar{z} \ln(\bar{z})}{1-r_\omega}. \quad (\text{D3}) \end{aligned}$$

Case $r_\omega > 1$. The density of plectonemes cannot be zero when $\bar{z} \downarrow 0$, since then l_p has to be 1 and the argument would go below $-1/e$ where the Lambert function is not real. So $\mu_z \rightarrow \infty$ and thus the argument of W_m goes to zero, from which it follows that $l_p \downarrow 0$ and consequently $\mu \uparrow 1$. Here we

find as asymptotic

$$\lim_{\bar{z} \rightarrow 0} l_p \simeq \frac{r_\omega}{r_\omega - 1} \bar{z}.$$

Case $r_\omega = 1$. Now the plectoneme number density goes to zero as $\mu \sim \sqrt{\bar{z}} e^{-\Delta'/2}$

Note that these two opposite limits at vanishing extension do not depend on Δ or its renormalized primed version.

These expressions are useful to obtain a parametrization of the μ , l_p , or \bar{z} curves as functions of r_n or v . Since

$$\mu = \frac{\mu_z \mu_l}{\mu_z \mu_l + \mu_z + \mu_l}, \quad r_n = \frac{(1 + r_\omega \mu_l) \mu_z}{\mu_z \mu_l + \mu_z + \mu_l}, \quad (\text{D4})$$

we can use this and Eq. (D2a) as a parametrization of the $\mu(r_n)$ curve by μ_z . At the start of the slope μ_z is zero. We just need to know its value at the end of the slope, which follows from the previous scaling relations as being infinite. From Eq. (D4) we then obtain that the maximum value r_n reaches

$$\lim_{\mu_z \rightarrow \infty} r_n = \begin{cases} 1 & \text{if } r_\omega \leq 1, \\ r_\omega & \text{if } r_\omega > 1, \end{cases} \quad (\text{D5})$$

which just tells us how much linking number the chain can absorb. More important is that if one wishes to interpret the turn-extension slopes in terms of a single plectoneme then when $r_\omega > 1$ the plectoneme parameters have to be adjusted. Some plots generated by this parametrization are in Fig. 4.

It is of practical importance to know when multiple plectonemes become significant not far after the transition, since it is there where most measurements are performed. We expect single-plectoneme behavior when most additional linking number goes into a growing plectoneme, or

$$l_p \simeq r_n \Rightarrow l_p \gg r_\omega \mu. \quad (\text{D6a})$$

From Eq. (D2a) and the properties of W_p it follows that close to the transition, where $\bar{z} \simeq 1$,

$$\frac{r_\omega}{\mu_z} e^{-\Delta'} \ll 1 \Rightarrow \mu \gg r_\omega e^{-\Delta'}. \quad (\text{D6b})$$

Finally, combining Eqs. (D6a) and (D6b) results in the following inequality that a mostly single-plectoneme configuration should obey:

$$l_p \gg \zeta := r_\omega^2 e^{-\Delta'}. \quad (\text{D6c})$$

Since close to the transition side of the turn-extension slope $l_p \ll 1$, the *multiplectoneme* factor ζ is an indicator for the appearance of several plectonemes. Once ζ becomes of the order 1, the change in the number of plectonemes plays an important part in the conversion of added linking number into writhe. Figure 5(a) shows the resulting ζ over a range of salt concentrations and forces.

The maximum of the loop density over the full range of allowed linking number densities is also straightforward to calculate to lowest order:

$$\mu_{\max} \simeq e^{-\Delta'/4} \frac{\cosh\left(\frac{\Delta'}{4}\right) - r_\omega e^{-\Delta'/4}}{\sinh\left(\frac{\Delta'}{4}\right) + (1 - r_\omega)r_\omega e^{-\Delta'/4}} \left[r_\omega e^{-\Delta'/4} + \left(1 - \sqrt{\frac{\sinh\left(\frac{\Delta'}{4}\right) + r_\omega e^{-\Delta'/4}}{\cosh\left(\frac{\Delta'}{4}\right) - r_\omega e^{-\Delta'/4}}}\right) \sinh\left(\frac{\Delta'}{4}\right) \right]. \quad (\text{D7})$$

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- [1] S. B. Smith, L. Finzi, and C. Bustamante, *Science* **258**, 1122 (1992).
 - [2] J. F. Marko and E. D. Siggia, *Macromolecules* **28**, 8759 (1995).
 - [3] T. Odijk, *Macromolecules* **28**, 7016 (1995).
 - [4] T. R. Strick, J.-F. Allemand, D. Bensimon, A. Bensimon, and V. Croquette, *Science* **271**, 1835 (1996).
 - [5] J. D. Moroz and P. Nelson, *Macromolecules* **31**, 6333 (1998).
 - [6] J. Gore, Z. Bryant, M. Nöllmann, M. Le, N. Cozzarelli, and C. Bustamante, *Nature (London)* **442**, 836 (2006).
 - [7] B. C. Daniels, S. Forth, M. Y. Sheinin, M. D. Wang, and J. P. Sethna, *Phys. Rev. E* **80**, 040901 (2009).
 - [8] T. Odijk, *J. Polym. Sci., Polym. Phys. Ed.* **15**, 477 (1977).
 - [9] J. Skolnick and M. Fixman, *Macromolecules* **10**, 944 (1977).
 - [10] G. S. Manning, *J. Chem. Phys.* **51**, 924 (1969).
 - [11] G. Călugăreanu, *Rev. Roum. Math. Pures Appl.* **4**, 257 (1959).
 - [12] J. White, *Am. J. Math.* **91**, 693 (1969).
 - [13] F. Fuller, *Proc. Natl. Acad. Sci. USA* **75**, 3557 (1978).
 - [14] J. Aldinger, I. Klapper, and M. Tabor, *J. Knot Theor. Ramif.* **4**, 343 (1995).
 - [15] M. Nizette and A. Goriely, *J. Math. Phys.* **40**, 2830 (1999).
 - [16] T. Odijk and A. Houwaart, *J. Polym. Sci., Polym. Phys. Ed.* **16**, 627 (1978).
 - [17] J. Ubbink and T. Odijk, *Biophys. J.* **76**, 2502 (1999).
 - [18] J. Philip and R. Wooding, *J. Chem. Phys.* **52**, 953 (1970).
 - [19] G. Téllez and E. Trizac, *J. Stat. Mech.: Theory Exp.* (2006) P06018.
 - [20] N. Clauvelin, B. Audoly, and S. Neukirch, *J. Comput.-Aided Mater. Des.* **14**, 95 (2007).
 - [21] P. K. Purohit, *J. Mech. Phys. Solids* **56**, 1715 (2008).
 - [22] C. Maffeo, R. Schöpfli, H. Brutzer, R. Stehr, A. Aksimentiev, G. Wedemann, and R. Seidel, *Phys. Rev. Lett.* **105**, 158101 (2010).
 - [23] S. Neukirch, *Phys. Rev. Lett.* **93**, 198107 (2004).
 - [24] S. Neukirch and J. F. Marko, *Phys. Rev. Lett.* **106**, 138104 (2011).
 - [25] M. Sheinin and M. Wang, *Phys. Chem. Chem. Phys.* **11**, 4800 (2009).
 - [26] J. F. Marko, *Phys. Rev. E* **57**, 2134 (1998).
 - [27] I. M. Kulić, H. Mohrbach, R. Thakkar, and H. Schiessel, *Phys. Rev. E* **75**, 011913 (2007).
 - [28] A. Cherstvy, *Eur. Biophys. J.* **40**, 69 (2011).
 - [29] T. Odijk and J. Ubbink, *Physica A* **252**, 61 (1998).
 - [30] M. Emanuel, *J. Chem. Phys.* **138**, 034903 (2013).
 - [31] A. Crut, D. A. Koster, R. Seidel, C. Wiggins, and N. H. Dekker, *Proc. Natl. Acad. Sci. USA* **104**, 11957 (2007).
 - [32] N. Hoskin and S. Levine, *Philos. Trans. R. Soc. London, Ser. A* **248**, 449 (1956).
 - [33] J. F. Marko and S. Neukirch, *Phys. Rev. E* **85**, 011908 (2012).
 - [34] H. Risken, *Fokker-Planck Equation* (Springer, Berlin, 1989).
 - [35] H. Brutzer, N. Luzzietti, D. Klaue, and R. Seidel, *Biophys. J.* **98**, 1267 (2010).
 - [36] J. F. Allemand, D. Bensimon, R. Lavery, and V. Croquette, *Proc. Natl. Acad. Sci. USA* **95**, 14152 (1998).
 - [37] F. Mosconi, J. Allemand, D. Bensimon, and V. Croquette, *arXiv:0809.0621*.
 - [38] T. Strick, J. Allemand, V. Croquette, and D. Bensimon, *Prog. Biophys. Mol. Biol.* **74**, 115 (2000).
 - [39] J. F. Marko, *Phys. Rev. E* **76**, 021926 (2007).
 - [40] C. Deufel, S. Forth, C. Simmons, S. Dejgosha, and M. Wang, *Nat. Methods* **4**, 223 (2007).
 - [41] S. Forth, C. Deufel, M. Y. Sheinin, B. Daniels, J. P. Sethna, and M. D. Wang, *Phys. Rev. Lett.* **100**, 148301 (2008).
 - [42] F. Mosconi, J. F. Allemand, D. Bensimon, and V. Croquette, *Phys. Rev. Lett.* **102**, 078301 (2009).
 - [43] M. van Loenhout, M. de Grunt, and C. Dekker, *Science* **338**, 94 (2012).
 - [44] S. Leikin, D. C. Rau, and V. A. Parsegian, *Phys. Rev. A* **44**, 5272 (1991).
 - [45] M. Emanuel (unpublished).
 - [46] T. Burkhardt, *J. Phys. A* **28**, L629 (1995).
 - [47] “NIST Digital Library of Mathematical Functions”, <http://dlmf.nist.gov/>, Release 1.0.5, online companion to [48].
 - [48] *NIST Handbook of Mathematical Functions*, edited by F. W. J. Olver, D. W. Lozier, R. F. Boisvert, and C. W. Clark (Cambridge University Press, New York, 2010), print companion to [47].