Inferring the effective thickness of polyelectrolytes from stretching measurements at various ionic strengths: applications to DNA and RNA

Ngo Minh Toan\textsuperscript{1,2} and Cristian Micheletti\textsuperscript{1}

\textsuperscript{1} International School for Advanced Studies (SISSA) and INFN, Via Beirut 2-4, 34014 Trieste, Italy
\textsuperscript{2} Institute of Physics, 10 Dao Tan, Hanoi, Vietnam

E-mail: ngo@sissa.it and michelet@sissa.it

Received 29 September 2005, in final form 16 January 2006
Published 24 March 2006
Online at stacks.iop.org/JPhysCM/18/S269

Abstract

By resorting to the thick-chain model we discuss how the stretching response of a polymer is influenced by the self-avoidance entailed by its finite thickness. The characterization of the force versus extension curve for a thick chain is carried out through extensive stochastic simulations. The computational results are captured by an analytic expression that is used to fit experimental stretching measurements carried out on DNA and single-stranded RNA (poly-U) in various solutions. This strategy allows us to infer the apparent diameter of two biologically relevant polyelectrolytes, namely DNA and poly-U, for different ionic strengths. Due to the very different degrees of flexibility of the two molecules, the results provide insight into how the apparent diameter is influenced by the interplay between the (solution-dependent) Debye screening length and the polymers’ ‘bare’ thickness. For DNA, the electrostatic contribution to the effective radius, $\Delta$, is found to be about five times larger than the Debye screening length, consistent with previous theoretical predictions for highly charged stiff rods. For the more flexible poly-U chains the electrostatic contribution to $\Delta$ is found to be significantly smaller than the Debye screening length.

1. Introduction

In recent years, the remarkable advancement of single-molecule manipulation techniques has made it possible to characterize with great accuracy how various biopolymers respond to mechanical stretching \cite{1–14}. The wealth of collected experimental data has constituted and still represents an invaluable and challenging benchmark for models of polymers’
elasticity [15–24]. The interpretation of single-molecule stretching experiments often relies on one-dimensional non-self-avoiding models of polymers. It is physically appealing that the schematic nature of such descriptions often conjugates with the capability of reproducing well experimental measurements. Two notable instances are represented by the freely jointed chain [25, 15] and the worm-like chain models which, in their original or extended forms, constitute the most commonly used theoretical frameworks for biopolymers’ stretching [26, 17, 27]. Both models are endowed with a parameter, the Kuhn length or the persistence length, that provides a phenomenological measure of the polymer stiffness and that is obtained by fitting the experimental data. It is important to notice, however, that it is possible to go beyond this phenomenological approach and connect the persistence length to the fundamental structural properties of a polymer. A strong indication of the feasibility of this scheme is provided by the fact that, for a large number of biopolymers, the observed persistence length shows an approximate quartic dependence on the polymer diameter, as predicted for ‘ideal’ elastic rods [15].

From this perspective it appears natural to investigate in detail the connection between structural properties and stretching response of biopolymers. We have recently pursued this objective by modelling explicitly the intrinsic thickness of a homo-polymer (treated as a tube with uniform cross-section) and characterizing the stretching response [21]. The theoretical and numerical results were employed in an appealing chemico-physical framework, where the diameter of a biopolymer was not probed directly but inferred through the mere knowledge of the stretching response. The adopted thick-chain model [28–30], briefly outlined in the next section, was used to fit stretching measurements obtained for a variety of biopolymers: DNA [5], the PEVK-domain of the titin protein [7, 8] and cellulose [31, 10]. For uncharged polymers, such as titin and cellulose, the effective diameters recovered from fitting the force-extension curves were very consistent with the stereochemical ones, thereby validating the thick-chain model approach [21]. Even more interesting is the case of polymers possessing a substantial linear charge density, such as DNA and RNA which will be the focus of the present study. The properties of polyelectrolytes, in fact, depend very strongly on the electrostatic screening provided by the ions present in solution. The influence of the electrostatic screening on the behaviour of polyelectrolytes has been extensively investigated both experimentally and theoretically [32–38]. From the latter perspective, it is customary to introduce apparent (or effective) physico–chemical parameters to describe the properties of a polyelectrolyte in a given solution with reference to the uncharged polymer case. For example, in the context of elasticity, one introduces an effective (solution-dependent) bending rigidity to account for the additional electrostatic contribution to the ‘bare’ persistence length of the hypothetically neutral polymer [37, 38]. Also, in the context of colloidal dispersions of stiff polyelectrolytes, one can describe the polymer as uncharged cylinders and resort to the theory for second virial coefficient to derive its solution-dependent effective diameter [34, 39, 40]. For DNA in solutions of low ionic strength, both the effective persistence length and effective diameter can exceed by several factors the bare ones. So far, these effective DNA properties have been typically probed by distinct methodologies. For example, stretching experiments were employed to establish the dependence of the persistence length on ionic strength [41, 6], while measurements of second virial coefficients, knotting probabilities or braiding properties were used for the effective diameter [42, 40, 43–45].

The thick-chain framework is used to obtain, starting from stretching measurement data, the effective diameter of a polyelectrolyte and to further relate it to its the effective persistence length. Besides the implications in the general context of polyelectrolytes the proposed method can be used to establish the effective structural parameters in coarse-grained studies of looping, knotting and packaging of biopolymers [46, 47].
Inferring the effective thickness of polyelectrolytes from stretching measurements

Non–local triplet

Local triplet

\[ \Delta \]

\[ \Delta \]

\[ \Delta \]

\[ \Delta \]

\[ \Delta \]

\[ \Delta \]

Figure 1. The finite thickness introduces steric constraints of local and non-local character that forbid configurations where the tube self-intersects. These constraints are conveniently treated within the three-body prescription of the thick-chain model. Within this approach the centreline of a viable configuration is such that the radii, \( r_{ijk} \), of the circles going through any triplet of points on the curve \( i, j, k \) are not smaller than \( \Delta \).

2. The thick-chain model

To characterize the stretching response of a polymer with finite thickness we shall view the latter as a tube with a uniform circular section in the plane perpendicular to the tube centreline. The chain thickness, \( \Delta \), is defined as the radius of the circular section. Several frameworks have been introduced to capture the uniform thickness constraint in a way apt for numerical implementation. These approaches typically rely on a discretized representation of the thick chain [48–50, 28, 51]. In this study we shall employ the piece-wise linear modelling of the chain centreline introduced by Gonzalez and Maddocks [28].

We shall indicate with \( \Gamma \) the centreline of the chain consisting of a succession of points \( \{\vec{r}_0, \vec{r}_1, \ldots\} \) equispaced at distance \( a \). We shall further denote with \( \vec{b}_i \) the virtual bond vector joining the \( i \)th and \( i+1 \)th points, \( \vec{b}_i = \vec{r}_{i+1} - \vec{r}_i \). In order for the succession of points \( \{\vec{r}_i\} \) to be a viable centreline for a chain of thickness \( \Delta \), it is necessary that the radii \( r_{ijk} \) of the circles going through any triples of points \( i, j, k \) are not smaller than \( \Delta \). Accordingly, the Hamiltonian for the thick-chain (tube) model can be written as

\[ H_{TC}(\Gamma) = \sum_{ijk} V_3(r_{ijk}) \tag{1} \]

where \( V_3 \) is the three-body potential used to enforce the thickness \( \Delta \) of the chain [28, 49, 52, 29, 30, 53]. As anticipated, the argument of \( V_3 \) is the radius of the circle going through the triplet of distinct points \( i, j, k \) and has the form

\[ V_3(r) = \begin{cases} 0 & \text{if } r > \Delta, \\ +\infty & \text{otherwise.} \end{cases} \tag{2} \]

Physically, the model of equation (1) introduces conformational restrictions for the centreline that are both local and non-local in character, as depicted in figure 1. The local constraints are those where the triplet \( i, j, k \) identifies three consecutive points. The limitation on the radius of the associated circumcircle reflects the fact that, to avoid singularities, the local radius of curvature must not be smaller than \( \Delta \). This reflects on the following bound on the angle formed by two consecutive bonds:

\[ \frac{\vec{b}_i \cdot \vec{b}_{i+1}}{a^2} \geq 1 - \frac{a^2}{2\Delta^2}. \tag{3} \]
On the other hand, there is also a non-local effect due to the fact that any two portions of the centreline at a finite arc-length separation cannot interpenetrate. It has been shown in [28] that this second effect can be accounted for by requiring that the minimum radius among circles going through any triplet of non-consecutive points is also greater than (or equal to) $\Delta$. The seamless way in which the local and non-local steric effects are accounted for makes the model particularly appealing. Other discrete models relying on pairwise interactions for the excluded volume (such as the cylindrical model of [44, 45]) may be adopted, though \textit{ad hoc} prescriptions for dealing with e.g. overlapping consecutive units need to be introduced [54].

In the present context we will consider the application of a stretching force, $\bar{f}$, to the ends of a chain $\Gamma = \Gamma_0, \ldots, \Gamma_N$ of thickness $\Delta$ (the contour length therefore being $L_c = N\alpha$). The Hamiltonian of equation (1) needs to be complemented with the stretching energy

$$\mathcal{H} = \sum_{ijk} V_3(r_{ijk}) - \bar{f} \cdot (\bar{r}_N - \bar{r}_0).$$

(4)

As customary we shall characterize the force dependence of the average normalized projection of the end-to-end distance, $\bar{r}_N - \bar{r}_0$, along the direction of applied force:

$$x = \frac{\langle (\bar{r}_N - \bar{r}_0) \cdot \bar{f} \rangle}{N\alpha |\bar{f}|}$$

(5)

where the brackets denote the canonical ensemble average. Owing to its self-avoiding nature, the stretching response of the chain cannot be characterized exactly by available analytical methods. We shall therefore resort to extensive Monte Carlo samplings, based on the Metropolis scheme, to evaluate the ensemble averages of equation (5).

Besides the numerical study of the tube model subject to stretching it is interesting to illustrate a simplification of the model of equation (1), which is amenable to an extensive analytical characterization. To do so we retain only the \textit{local} thickness constraint and thus end up with a model that is essentially non-self-avoiding. The simplified nature of this problem, however, makes it very tractable also in the presence of a bending rigidity penalty, $\kappa_b$. We shall therefore consider the Hamiltonian

$$\mathcal{H} = \sum_i V_3(r_{i,i+1,i+2}) - \bar{f} \cdot (\bar{r}_N - \bar{r}_0) - \kappa_b \sum_i \frac{\bar{b}_i \cdot \bar{b}_{i+1}}{a^2},$$

(6)

again we stress that the three-body potential is restricted only to consecutive (local) triplets. In this form the stretching response of the model can be characterized exactly both at very low and very high forces using standard statistical–mechanical procedures [55, 56]. These two limiting regimes can be joined together to yield the following approximate expression for the stretching response of a locally thick chain with bending rigidity (LTC + BR, for brevity):

$$\beta a f = 2K \left[ 1 + \left( \frac{1}{2K} \right)^2 \frac{1}{(1-x)^2} - 1 + \left( \frac{1}{2K} \right)^2 \right] + \frac{3}{1 + y(K, \Delta/a)} \frac{1}{2K \sqrt{1 + (1/2K)^2}} x$$

(7)

where $K = \beta \kappa_b$, $\beta = 1/K_B T$ is the inverse Boltzmann factor and

$$y(K, \Delta/a) = \begin{cases} 1 - \frac{a^2}{2\Delta^2} \left( \frac{1}{1 - e^c} + \frac{1}{c} \right) & \Delta/a > 0.5 \\ \coth(K) - \frac{1}{K} & \Delta/a \leq 0.5 \end{cases}$$

(8)
Inferring the effective thickness of polyelectrolytes from stretching measurements

with \( z = \frac{a^2}{\Delta} K \). The two cases in the above equation reflect the fact that, for \( \Delta < a/2 \), no restriction applies to the angle formed by two consecutive virtual bonds since inequality (3) is always satisfied.

Expression (7) possesses some noteworthy limits. First, in the absence of thickness and in the continuum limit \( (a \to 0, K \to \xi_p/a, \xi_p \) being the persistence length) \([55, 56]\), the model reduces to the well known Marko and Siggia result for the WLC:

\[
f(x) = \frac{k_B T}{\xi_p} \left[ \frac{1}{4(1 - x)^2} - \frac{1}{4} + x \right].
\] (9)

Second, in the absence of both thickness and bending rigidity one recovers the low- and high-force response of the freely jointed chain with Kuhn length equal to \( a \):

\[
f(x) \approx \begin{cases} 
3k_B T, & x \to 0; \\
\frac{a}{1}, & x \to 1.
\end{cases}
\] (10)

Also of interest is the case of finite thickness \( \Delta/a > 0.5 \) but no bending rigidity. In this case one obtains the following expression for the persistence length:

\[
\xi_p = -\frac{a}{\ln \left( 1 - \frac{a^2}{4\Delta} \right)}.
\] (11)

Though this expression does not include the non-local self-avoiding effects it will be shown later to provide a good approximation of the persistence length obtained numerically for the full model of equation (1).

We conclude this section by emphasizing that for the models of equations (1) and (6) the spacing of consecutive points is constant along the centreline, so that the contour length is unaffected by the application of forces of arbitrary strength. The inextensibility property is obviously a simplification of the behaviour found in naturally occurring polymers which, at very high forces, can undergo isomerization or structural transitions resulting in an ‘overstretching’ beyond their nominal contour length. Several approximate treatments have been developed to correct the stretching response of inextensible models so to account for overstretching \([57, 5]\) by adopting additional parameters in the theory. In the present study we shall keep the number of model parameters to a minimum and hence postpone to future work the investigation of the most suitable way to include overstretching in the TC model.

3. Numerical results

The characterization of the stretching response of the thick chain of equation (4) was carried out using a Monte Carlo scheme: starting from an arbitrary initial chain configuration satisfying the thickness constraints, the exploration of the available structure space was done by distorting conformations by means of pivot and crankshaft moves. Newly generated structures are accepted/rejected according to the standard Metropolis criterion (the infinite strength of the three-body penalties of equation (2) was enforced by always rejecting configurations violating the circumradius constraints).

The discretization length, \( a \), was taken as the unit length in the problem and several values of \( \Delta/a \) were considered, ranging from the minimum allowed value of 0.5 to the value of 4.0. This upper limit appears adequate in the present context since the largest nominal ratio for \( \Delta/a \) among the biopolymers considered here is achieved for dsDNA for which one has \( \Delta/a \approx 3.7 \) \([47]\). For each explored value of \( \Delta/a \) considered, we considered chains of
length at least ten times bigger than the persistence length estimated through equation (11). The relative elongation of the chain was calculated for increasing values of the applied stretching force (typically about 100 distinct force values were considered). For each run, after equilibration, we measured the autocorrelation time and sampled a sufficient number of independent conformations to achieve a relative error of, at most, $10^{-3}$ on the average chain elongation. For moderate or high forces this typically entailed the collection of $10^4$ independent structures while a tenfold increase of sampling was required at small forces due to the broad distribution of the end-to-end separation along the force direction.

We first discuss the results for the persistence length obtained from the decay of the tangent–tangent correlations measured at zero force over an ensemble of sampled structures picked at times greater than the system autocorrelation time. The resulting data are shown in figure 2(a), along with the curve corresponding to the approximate expression of equation (11). It can be seen that the local approximation for the persistence length is very good in the range $1.0 \leq \Delta \leq 4$, where the relative difference from the value found numerically is typically inferior to 10%. Significant relative discrepancies are, instead, found as $\Delta$ approaches the limiting value of 0.5 (though it should be noted that the single-exponential fit suffers from the very rapid decay of the tangent autocorrelation). In this case, only a narrow range of values for the angle formed by two consecutive bonds is forbidden. Consequently, the persistence length is very much affected by the (non-local) self-avoidance condition that is unaccounted for by the simple expression of equation (11). As intuitively expected the value of $\xi_p$ found numerically is larger than the one based on the local-thickness approximation.

The analysis of the numerical results revealed several different stretching regimes in the elastic response of a thick chain. These are best discussed in connection with analogous regimes found in the two standard reference models, the freely jointed chain (FJC) and the worm-like chain (WLC). We first point out that for both these models, as well as the generalization of equation (7), the relative elongation, $x$, depends linearly on the applied force, $f$. This result also holds for the thick-chain model. However, due to the inclusion of the self-avoidance effect in the TC (absent in both the FJC and WLC), the Hookean relationship between $f$ and $x$ disappears in the limit of long polymer chains in favour of the Pincus regime, $f \sim x^{1/(1/v-1)}$. 

---

**Figure 2.** (a) Comparison between the persistence length of a thick chain obtained by Monte Carlo sampling (dotted–squares) and that obtained from the local thickness approximation (solid curve) of equation (11). The inset illustrates the limitations of the local approximation for low values of $\Delta/a$.

(b) Tangent–tangent correlation (dotted–squares) obtained from the analysis of $10^4$ (uncorrelated) Monte Carlo-generated chains of 1000 segments and with $\Delta/a = 2$. The error bars which show the uncertainty of the plotted values are smaller than the square symbols. The solid curve is a single-exponential fit to the numerical data, yielding $\xi_p = 16.71 \pm 0.03$. 

Figure 3. Linear (a) and semi-log (b) plots of the force versus relative extension curves obtained from Monte Carlo sampling of chains of relative thickness $\Delta/a = 0.75, 1.0, 2.0$ and 3.

Figure 4. (a) Extension versus reduced force for a chain of relative thickness $\Delta/a = 1.0$. Data points are presented in the $(\log(1-x), \log(f))$ plane to highlight the WLC- and FJC-like regimes found at moderate and high forces. (b) Illustration of the low-force crossover from the Hookean regime, $x \propto f$, to the Pincus one, $x \propto f^{2/3}$ for a chain of 1200 segments and $\Delta/a = 1.0$.

$\nu \sim 3/5$ being the critical exponent for self-avoiding polymers in three dimensions [25, 16, 58]; see figure 4.

For intermediate forces the Pincus behaviour is found to be followed by a second regime characterized by the same scaling relation as found in the WLC at high forces, $f \sim (1-x)^{-2}$. As shown in figure 4, at still higher forces the same scaling relation as found in the FJC is observed, $f \sim (1-x)^{-1}$. Physically, the first two regimes are determined by self-avoidance and chain stiffness or persistence length while the last regime is ascribable to the discrete nature of the chain [59, 60, 55, 61].

In order to apply the thick-chain model to contexts where experimental data are available, we have analysed the data of the numerical simulations with the purpose of extracting an analytical expression capturing the observed functional dependence of $f$ on $\Delta$ and $a$. For any value of $a$ and $\Delta$ the sought expression should reproduce the succession of the three regimes discussed above. Among several trial formulae we found, a posteriori, that the best interpolation was provided by the expression

$$f(x) = \frac{k_B T}{a(1-x)} \tanh \left( \frac{k_1 x^{3/2} + k_2 x^2 + k_3 x^3}{1-x} \right).$$

(12)
Figure 5. Illustration of the performance of the parametric expression of equation (12) in reproducing the stretching response obtained numerically for chains of relative thickness $\Delta/a = 0.75$ and $\Delta/a = 4.0$ ((a) and (b), respectively). In both cases the discrepancy between the computed and parametrized values is about 1% on average (and always smaller than 5%).

where the parametric dependence on $\Delta$ and $a$ is carried by the following expressions for the $k$:

\[
\begin{align*}
  k_1^{-1} &= -0.28394 + 0.76441 \Delta/a + 0.31858 \Delta^2/a^2, \\
  k_2^{-1} &= +0.15989 - 0.50503 \Delta/a - 0.20636 \Delta^2/a^2, \\
  k_3^{-1} &= -0.34984 + 1.23330 \Delta/a + 0.58697 \Delta^2/a^2.
\end{align*}
\]  

Within the explored ranges of $\Delta$ and $a$, the relative extension obtained from equation (12) differs on average by 1% (and at most by 5%) from the true values at any value of the applied force, as shown in figure 5.

4. Applications to experimental data and discussions

We shall now discuss the application of the TC model to data sets obtained in DNA and RNA stretching experiments carried out for various $[\text{Na}^+]$ concentrations. In particular, the data for dsDNA are taken from [2] (solutions of 10, 1.0 and 0.1 mM $[\text{Na}^+]$) while for single-stranded RNA (poly-U) we considered the recent results of [62] (solutions of 500, 300, 100, 50, 10 and 5 mM $[\text{Na}^+]$). The fits of the experimental data are carried out through the standard procedure of minimizing the $\chi^2$ over the TC parameters: the contour length, $L_c$, chain granularity, $a$ and chain thickness, $\Delta$. For the calculation of $\chi^2$ we have estimated the effective uncertainty on the force measurements, taking the relative extension as the independent variable, directly from the large data sets of poly-U (more than 400 points for each set). For dsDNA, owing to the more limited number of points (about 20–25) we propagated the declared experimental uncertainty on extension.

The results of the fit procedures are provided in graphs 6 and 7 and tables 1 and 2. It is particularly appealing that, over the about 400 data points available for poly-U, the $\chi^2$ associated with the thick chain is very close to unity for the set of measurements carried out for $[\text{Na}^+]$ in the 50–500 mM range. In the case of the two smallest concentrations, $[\text{Na}^+] = 5$ and 10 mM, a significant increase of the $\chi^2$ is observed. The same is true for the fit of DNA measurements carried out in 0.1 mM $[\text{Na}^+]$. The worsening of the TC performance consequent to the increase of the screening length is reflected, among other effects, by the progressive importance of accounting for overstretching [63, 6, 62]. Accordingly, the fitting parameters obtained at the lowest salt concentrations can be expected to change upon the introduction of a stretching modulus.
Inferring the effective thickness of polyelectrolytes from stretching measurements

Figure 6. Thick-chain fit of the experimental data (squares, circles and triangles) on dsDNA in solutions of various ionic strengths. The best-fit parameters are provided in table 1.

Figure 7. Application of the thick-chain model to poly-U stretching data for (a) 300 mM [Na\(^{+}\)] and (b) 10 mM [Na\(^{+}\)]. Experimental data are shown as open circles; the fit with the TC is denoted with a solid line. The best-fit parameters for the TC are provided in table 2.

Table 1. Best-fit parameters obtained by applying the parametric force–extension expressions of the TC model to the experimental data on DNA in solutions of various ionic strengths.

| dsDNA | [Na\(^{+}\)] (mM) | \(\alpha\) (nm) | \(\Delta\) (nm) | \(\xi_p\) (nm) | \(L_c\) (nm) | \(\chi^2\) |
|-------|-----------------|----------------|----------------|---------------|-------------|---------|
|       |                 |                |                |               |             |         |
| 10    | 9 ± 2           | 12 ± 2         | 55 ± 2         | 32.9 ± 0.2    | 0.6         |
| 1.0   | 25 ± 7          | 26 ± 4         | 94 ± 4         | 32.3 ± 0.2    | 0.3         |
| 0.1   | 36 ± 16         | 48 ± 13        | 242 ± 14       | 32.3 ± 0.2    | 1.9         |

For both DNA and RNA the viability of the structural and elastic parameters of the TC fit can be compared against those obtained by different models and physical approaches. The most natural term of comparison for the elastic response is provided by the widely employed WLC, which is the common reference model for determining the persistence length and contour length of several types of biopolymers. It is appealing that both \(\xi_p\) and \(L_c\) given by the TC for dsDNA and poly-U in tables 1 and 2 are in close agreement with the same quantities given by the WLC, the relative difference being typically less than 5\% for both quantities. For dsDNA, the values
Table 2. Best-fit parameters obtained by applying the parametric force–extension expressions of
the TC model to the experimental data on poly-U in solutions of various ionic strengths.

| [Na\(^+\)] (mM) | \(a\) (nm) | \(\Delta\) (nm) | \(\xi_p\) (nm) | \(L_c\) (nm) | \(\chi^2\) |
|-----------------|----------|-------------|-------------|-------------|---------|
| 500             | 1.10 ± 0.01 | 0.64 ± 0.01 | 0.80 ± 0.01 | 2126 ± 4    | 1.49    |
| 300             | 1.09 ± 0.02 | 0.65 ± 0.01 | 0.91 ± 0.01 | 2123 ± 5    | 1.29    |
| 100             | 1.08 ± 0.03 | 0.68 ± 0.02 | 1.11 ± 0.01 | 2117 ± 8    | 1.05    |
| 50              | 1.11 ± 0.03 | 0.73 ± 0.02 | 1.30 ± 0.01 | 2134 ± 8    | 1.33    |
| 10              | 1.33 ± 0.08 | 0.94 ± 0.04 | 1.94 ± 0.03 | 2138 ± 12   | 4.21    |
| 5               | 1.41 ± 0.10 | 1.03 ± 0.05 | 2.26 ± 0.05 | 2142 ± 12   | 4.34    |
Inferring the effective thickness of polyelectrolytes from stretching measurements

Figure 8. Effective radius, $\Delta$, of poly-U as a function of ionic strengths. The dashed curve is a visual guideline obtained from a spline interpolation of the points obtained from the TC fits. The approximate linear dependence of $\Delta$ on $1/\sqrt{[Na^+]}$ (with $[Na^+]$ expressed in molar units) is shown in the inset.

Theoretical results derived by Stigter and Odijk [34, 39] have been previously used to predict the effective diameter of double-stranded DNA. The available theoretical predictions based on the approach of [34, 40, 39] yield dsDNA radii of 8.1 nm for 10 mM $[Na^+]$, 27.1 nm for 1 mM $[Na^+]$ and 96.3 for 0.1 mM $[Na^+]$. Given the very diverse nature of the approach and approximations of the present study and of [34, 39], it is pleasing that the two determinations of the effective thickness are in reasonable agreement.

We finally turn to the case of poly-U. Owing to its high flexibility, even short stretches of RNA cannot be modelled as stiff rods, and hence the theoretical analysis of [34] cannot be used to generate a comparison term for the effective radius, $\Delta$, of poly-U. It was pointed out by Stigter that for DNA the electrostatic contribution to the effective radius, though exceeding by several times the bare one, was of the same order as the Debye screening length, $\lambda_D$ (the ‘proportionality factor’, ranging from about four to eight) [34]. Also in the case of RNA, the Debye length, which is proportional to $1/\sqrt{[Na^+]}$, is a useful concept for rationalizing the dependence of the effective radius on the ionic strength. In fact, within the explored range of $[Na^+]$, $\Delta$ appears to increase linearly with $1/\sqrt{[Na^+]}$, as visible in the inset of figure 8(b).

Assuming the validity of such a linear relationship, the ‘bare’ radius of poly-U is estimated by extrapolating $\Delta$ for vanishing $\lambda_D$. One obtains $\Delta_{\text{bare}} \approx 0.59 \pm 0.01$ nm, which compares well with the nominal value of about 0.5 nm observed in crystallographic structures of (non-hydrated) poly-U fragments (from PDB structure 1I5L). Incidentally, we also mention that the value of $a$ agrees with the nominal size of the modular poly-U units, which is about 0.6 nm. Several efforts have been devoted in the past to establishing how the apparent persistence length of polyelectrolytes is affected by the interplay between the bare persistence length, the linear charge density and the ionic strength [37, 36, 38]. Different manners of functional dependence of $\xi_p$ on $\lambda_D$ are, in fact, observed for polyelectrolytes with different degrees of flexibility, such as double-stranded and single-stranded DNA. It is, therefore, interesting to compare the dependence of the effective diameter of DNA and poly-U. The most notable difference is that the proportionality factor between the electrostatic contribution to the effective radius and the Debye length is of the order of 0.1, therefore appreciably smaller than for the case of dsDNA.
5. Conclusions

Extensive stochastic sampling techniques were used to characterize the behaviour of an inextensible thick polymer subject to a stretching force. The extension versus force response found numerically was parametrized in terms of the polymer structural parameters (thickness and monomer length) and captured by an analytic expression. The resulting formula was used to fit experimental data obtained from stretching measurements of DNA and poly-U in solutions of different ionic strength. This represents a novel and physically appealing route to extract the apparent structural parameters of polyelectrolytes starting merely from their elastic response. The inferred effective diameter for DNA was found to be in satisfactory agreement with the estimates obtained by Stigter and Odijk through unrelated approaches. In particular, the electrostatic contribution to the effective DNA radius was found to be several times larger than the Debye screening length, $\lambda_D$. Moreover, for the much more flexible poly-U chain it is observed that the effective radius strongly depends on the ionic strength, having an approximately linear dependence on $\lambda_D$ within the available ranges of $[Na^+]$. At variance with dsDNA, however, the electrostatic contribution to the effective radius was found to be an order of magnitude smaller than the Debye screening length.

Acknowledgments

We are indebted to K Visscher for having provided us with the experimental data on poly-U and for useful comments. We thank T Odijk for valuable suggestions and are grateful to D Marenduzzo for several discussions and the careful reading of the manuscript.

References

[1] Bustamante C, Bryant Z and Smith S B 2003 Nature 421 423–7
[2] Smith S B, Finzi I and Bustamante C 1992 Science 258 1122–6
[3] Cluzel P, Lebrun A, Heller C, Lavery R, Visvoy J-L, Chatenay D and Caron F 1996 Science 271 792
[4] Strick T R, Allemand J F, Bensimon D, Bensimon A and Croquette V 1996 Science 271 1835–7
[5] Bouchiat C, Wang M D, Allemand J F, Strick T, Block S M and Croquette V 1999 Biophys. J. 76 409–13
[6] Wenner J R, Williams M C, Rouzina I and Bloomfield V A 2002 Biophys. J. 82 3160
[7] Linke W A, Kilke M, Li H, F-Becker S, Neagoe C, Manstein D J, Gautel M and Fernandez J M 2002 J. Struct. Biol. 137 194–205
[8] Li H, Oberhauser A F, Redick S D, C-Vazquez M, Erickson H P and Fernandez J M 2001 Proc. Natl Acad. Sci. USA 98 10682–6
[9] Rief M, Pascual J, Saraste M and Gaub H E 1999 J. Mol. Biol. 286 553–61
[10] Marszalek P E, Oberhauser A F, Pang Y-P and Fernandez J M 1998 Nature 396 661–4
[11] Williams P M, Fowler S B, Best R B, Toca-Herrera J L, Scott K A, Steward A and Clarke J 2003 Nature 422 446–9
[12] Best R B, Fowler S B, Toca-Herrera J L, Steward A, Paci E and Clarke J 2003 J. Mol. Biol. 330 867–77
[13] Wang M D, Yin H, Landick R, Gelens J and Block S M 1997 Biophys. J. 72 1335–46
[14] Strick T, Allemand J F, Croquette V and Bensimon D 2000 Prog. Biophys. Mol. Biol. 74 115–40
[15] Boal D 2002 Mechanics of the Cell (Cambridge: Cambridge University Press)
[16] Pincus P 1976 Macromolecules 9 386–8
[17] Bustamante C, Marko J F, Siggia E D and Smith S 1994 Science 265 1599–600
[18] Cieplak M, Hoang T X and Robbins M O 2002 Proteins 49 114–24
[19] Gao M, Wilmanss M and Schulten K 2002 Biophys. J. 83 3435–45
[20] Bensimon D, Dohmi D and Mezard M 1998 Europhys. lett. 42 97102
[21] Toan N M, Marenduzzo D and Micheletti C 2005 Biophys. J. 89 80–6
[22] Hamed S A 2005 Phys. Rev. E 72 (3) (Preprint cond mat/0409492)
[23] Hugel T, Rief M, Seitz M, Gaub H E and Netz R R 2005 Phys. Rev. Lett. 94 048301
[24] Piana S 2005 Nucleic Acids Res. 33 7029–38
Inferring the effective thickness of polyelectrolytes from stretching measurements

[25] Flory P J 1989 Statistical Mechanics of Chain Molecules (Munich: Hanser)
[26] Marko J F and Siggia E D 1995 Macromolecules 28 8759–70
[27] Hansen P L and Podgornik R 2001 J. Chem. Phys. 114 8637–48
[28] Gonzalez O and Maddocks J H 1999 Proc. Natl Acad. Sci. USA 96 4769–73
[29] Maritan A, Micheletti C, Trovato A and Banavar J R 2000 Nature 406 287
[30] Stasiak A and Maddocks J H 2000 Nature 406 251–3
[31] Marszalek P E, Pang Y-P, Li H, Yazal J E, Oberhauser A F and Fernandez J M 1999 Proc. Natl Acad. Sci. USA 96 7894–8
[32] Odijk T 1977 J. Polym. Sci. Polym. Phys. 15 477–83
[33] Skolnick J and Fixman M 1977 Macromolecules 10 944–8
[34] Sottas P-E, Larquet E, Stasiak A and Dubochet J 1999 Biophys. J. 77 1858–70
[35] Rybenkov V V, Cozzarelli N R and Vologodskii A V S 1993 Proc. Natl Acad. Sci. USA 90 5307