Dynamics of supercoiled DNA with complex knots: large-scale rearrangements and persistent multi-strand interlocking

Lucia Coronel¹, Antonio Suma¹,² and Cristian Micheletti¹,∗

¹SISSA - Scuola Internazionale Superiore di Studi Avanzati, Via Bonomea 265, 34136 Trieste, Italy and ²Institute for Computational Molecular Science, College of Science and Technology, Temple University, Philadelphia, PA 19122, USA

Received March 08, 2018; Revised May 22, 2018; Editorial Decision May 23, 2018; Accepted May 24, 2018

ABSTRACT

Knots and supercoiling are both introduced in bacterial plasmids by catalytic processes involving DNA strand passages. While the effects on plasmid organization has been extensively studied for knotting and supercoiling taken separately, much less is known about their concurrent action. Here, we use molecular dynamics simulations and oxDNA, an accurate mesoscopic DNA model, to study the kinetic and metric changes introduced by complex (five-crossing) knots and supercoiling in 2 kbp-long DNA rings. We find several unexpected results. First, the conformational ensemble is dominated by two distinct states, differing in branchedness and knot size. Secondly, fluctuations between these states are as fast as the metric relaxation of unknotted rings. In spite of this, certain boundaries of knotted and plectonemically-wound regions can persist over much longer timescales. These pinned regions involve multiple strands that are interlocked by the cooperative action of topological and supercoiling constraints. Their long-lived character may be relevant for the simplifying action of topoisomerases.

INTRODUCTION

The structural organization of bacterial plasmids is profoundly affected by homeostatic catalytic processes involving DNA such as transcription, replication and recombination. The best known example is the level of negative supercoiling, ~−5%, that is maintained by topoisomerases (1–3) and that creates an interplay between DNA twist (local winding around the double-helix axis) and writhe (winding of the double-helix axis around itself). Inter- and intramolecular strand passages catalysed by topoisomerases can affect DNA entanglement too. The specific types of knots and links that can be introduced (4–8) or removed (8–10) by these mechanisms, can vary with experimental conditions, the specific topoisomerase action on well-defined patterns of crossings (4) as well as DNA length (5–8,10–12), which is a key determinant of knot complexity in viral (13–16) and eukaryotic DNA too (17).

The structural constraints associated to supercoiling and knotting have functional implications, too. Plectonemes, for instance, control the degree of branchedness of DNA rings (18–21) and this, in turn, affects the contact probability of loci at large genomic separations (18,22–25). In addition, the mechanics of superhelical stress, which can have long-range effects (26,27), can facilitate the unzipping of AT-rich regions that are upstream of genes, and hence the binding of promoters (26,28–33).

Knots, on the other hand, have been mostly associated with detrimental functional effects, such as stalling DNA replication and transcription (5,6,34). In fact, their statistically inevitable emergence is counteracted by specific topoisomerases that can eventually remove them (10,35,36) via selective strand passages, arguably at hooked juxtapositions that are characteristic of knots (37–40).

Intriguingly, recent modelling studies by Stasiak’s lab, have pointed to a primary role of supercoiling in the removal of DNA knots, too (41–43). Stochastic simulations of coarse-grained DNA filaments tied in trefoil knots, the simplest non-trivial topology, have shown that the systematic accumulation of twist leads to a tightening, or localization, of the knot. This localization expectedly facilitates the local action (recognition and strand passage) of topoisomerases irrespective of knot chirality (41,42,44).

These results add a novel appealing layer to the functional role of supercoiling. At the same time, its interplay with knots, especially the more complex ones reported in plasmids, is largely unexplored and key questions are still unanswered. For instance: how do knots more complex than trefoils affect the branchedness of supercoiled rings? Would the latter be increased by the large writhe of complex knots, or would it be suppressed by the topological constraints? Also, what is the effect of an intricate topology on
the internal dynamics of plasmids, and how does it differ from the one of supercoiling? Do knots trap the system in long-lived states and, if so, what are their characteristics?

Here, we tackle these questions for 2 kbp-long DNA rings using molecular dynamics simulations and oxDNA (45–47), a model with an accurate mesoscopic representation of nucleotides and their interactions. This level of detail makes it possible to gather multi-ms trajectories for kbp-long-lived states and, if so, what are their characteristics? from the one of supercoiling? Do knots trap the system in the internal dynamics of plasmids, and how does it differ from the one of supercoiling? Do knots trap the system in long-lived states and, if so, what are their characteristics?

We considered 2 kbp-long DNA rings with or without negative supercoiling and in three different topologies: $5_1$ and $5_2$ left-handed knots as well as the unknot ($0_1$ topology).

The initial conformations were generated with the following two-tier scheme. First, we produced the centerline of the double-stranded DNA rings by using the KnotPlot software (available at www.knotplot.com) to create smooth, symmetric forms of $0_1$, $5_1$ and $5_2$ knots, see Figure 1A. To be consistent with experimental observations on pBR322 plasmids (5,16) the chirality of the five-crossing knots was set to be left-handed, i.e. projected crossings have negative sign.

These circular centerlines, discretised in 2000 segments, were next turned into the oxDNA double-helical representation by a fine-graining procedure, where each segment was mapped into the six interaction centers of the two paired nucleotides (48), see inset in Figure 1A. In this fine-graining procedure, the average twist between consecutive bases was adjusted differently for each topology to yield the sought level of supercoiling, as discussed in detail in refs. (42,50). Such twist adjustment is needed to account for the topological contribution to the writhe, i.e. the winding of the DNA centerline on itself. In fact, the canonical ensemble average of the writhe is usually different from zero for torsionally-relaxed knotted rings (51). For each considered topological state we accordingly adjusted the twist uniformly to deal with the two different cases of torsionally-relaxed DNA rings and negatively supercoiled ones. In the latter case we set the relative amount of supercoiling equal to $-0.05$, the typical homeostatic level in bacterial plasmids.

### Materials and Methods

#### Model

We considered 2 kbp-long DNA rings with or without negative supercoiling and in three different topologies: $5_1$ and $5_2$ left-handed knots as well as the unknot ($0_1$ topology).

The rings were described with the oxDNA mesoscopic model (45–47), where each nucleotide is represented by three interaction centers (after the sugar, base and phosphate groups) with average (sequence-independent) parameters for base pairing, coaxial stacking, steric and electrostatic interactions. For the latter we used the default oxDNA parameterization for 1M monovalent salt.

#### Initial setup

The initial conformations were generated with the following two-tier scheme. First, we produced the centerline of the double-stranded DNA rings by using the KnotPlot software (available at www.knotplot.com) to create smooth, symmetric forms of $0_1$, $5_1$ and $5_2$ knots, see Figure 1A. To be consistent with experimental observations on pBR322 plasmids (5,16) the chirality of the five-crossing knots was set to be left-handed, i.e. projected crossings have negative sign.

These circular centerlines, discretised in 2000 segments, were next turned into the oxDNA double-helical representation by a fine-graining procedure, where each segment was mapped into the six interaction centers of the two paired nucleotides (48), see inset in Figure 1A. In this fine-graining procedure, the average twist between consecutive bases was adjusted differently for each topology to yield the sought level of supercoiling, as discussed in detail in refs. (42,50). Such twist adjustment is needed to account for the topological contribution to the writhe, i.e. the winding of the DNA centerline on itself. In fact, the canonical ensemble average of the writhe is usually different from zero for torsionally-relaxed knotted rings (51). For each considered topological state we accordingly adjusted the twist uniformly to deal with the two different cases of torsionally-relaxed DNA rings and negatively supercoiled ones. In the latter case we set the relative amount of supercoiling equal to $-0.05$, the typical homeostatic level in bacterial plasmids.
We first constructed a contact map for the DNA centreline using a tolerant cutoff distance of 40 \( \text{bp} \). The latter two are left-handed, i.e. the topological sign of their projected crossings is negative, as indicated. The \( 5_1 \) and \( 5_2 \) snapshots have been edited to highlight the overlap of negative supercoiling (-5\%).

Detection of plectonemes

For each band we identified its apex, which is the site, \( i \), associated to the longest uninterrupted ladder of contacts \( \{i + \Delta, i - \Delta\}, (i + \Delta + 1, i - \Delta - 1), (i + \Delta + 2, i - \Delta - 2) \), starting at a sequence separation, \( \Delta \), not larger than 400 \text{ bp} to avoid excessively large apical loops. At the same time, trivial contacts due to short sequence separations \( \Delta < 100 \text{ bp} \) are not counted. We note that at this stage, and at the next one too, there could be overlaps in the regions covered by different putative plectonemes.

Within each band with \( \Delta < 400 \text{ bp} \), we then searched for the presence of tight contacts, a signature feature of superhelices. Specifically, we searched for the contacts at distance smaller than 7.5\( \sigma \) (\( \sim 6 \text{ Å} \)), and took the contact map diagonal. These bands correspond to dsDNA stretches in spatial proximity and with opposite directionality (in an oriented ring).

For each band we identified its apex, which is the site, \( i \), associated to the longest uninterrupted ladder of contacts \( \{i + \Delta, i - \Delta\}, (i + \Delta + 1, i - \Delta - 1), (i + \Delta + 2, i - \Delta - 2) \), starting at a sequence separation, \( \Delta \), not larger than 400 \text{ bp} to avoid excessively large apical loops. At the same time, trivial contacts due to short sequence separations \( \Delta < 100 \text{ bp} \) are not counted. We note that at this stage, and at the next one too, there could be overlaps in the regions covered by different putative plectonemes.

Within each band with \( \Delta < 400 \text{ bp} \), we then searched for the presence of tight contacts, a signature feature of superhelices. Specifically, we searched for the contacts at distance smaller than 7.5\( \sigma \) (\( \sim 6 \text{ Å} \)), and took the contact map diagonal. These bands correspond to dsDNA stretches in spatial proximity and with opposite directionality (in an oriented ring).

For each band we identified its apex, which is the site, \( i \), associated to the longest uninterrupted ladder of contacts \( \{i + \Delta, i - \Delta\}, (i + \Delta + 1, i - \Delta - 1), (i + \Delta + 2, i - \Delta - 2) \), starting at a sequence separation, \( \Delta \), not larger than 400 \text{ bp} to avoid excessively large apical loops. At the same time, trivial contacts due to short sequence separations \( \Delta < 100 \text{ bp} \) are not counted. We note that at this stage, and at the next one too, there could be overlaps in the regions covered by different putative plectonemes.

Within each band with \( \Delta < 400 \text{ bp} \), we then searched for the presence of tight contacts, a signature feature of superhelices. Specifically, we searched for the contacts at distance smaller than 7.5\( \sigma \) (\( \sim 6 \text{ Å} \)), and took the contact map diagonal. These bands correspond to dsDNA stretches in spatial proximity and with opposite directionality (in an oriented ring).

For each band we identified its apex, which is the site, \( i \), associated to the longest uninterrupted ladder of contacts \( \{i + \Delta, i - \Delta\}, (i + \Delta + 1, i - \Delta - 1), (i + \Delta + 2, i - \Delta - 2) \), starting at a sequence separation, \( \Delta \), not larger than 400 \text{ bp} to avoid excessively large apical loops. At the same time, trivial contacts due to short sequence separations \( \Delta < 100 \text{ bp} \) are not counted. We note that at this stage, and at the next one too, there could be overlaps in the regions covered by different putative plectonemes.

Within each band with \( \Delta < 400 \text{ bp} \), we then searched for the presence of tight contacts, a signature feature of superhelices. Specifically, we searched for the contacts at distance smaller than 7.5\( \sigma \) (\( \sim 6 \text{ Å} \)), and took the contact map diagonal. These bands correspond to dsDNA stretches in spatial proximity and with opposite directionality (in an oriented ring).

For each band we identified its apex, which is the site, \( i \), associated to the longest uninterrupted ladder of contacts \( \{i + \Delta, i - \Delta\}, (i + \Delta + 1, i - \Delta - 1), (i + \Delta + 2, i - \Delta - 2) \), starting at a sequence separation, \( \Delta \), not larger than 400 \text{ bp} to avoid excessively large apical loops. At the same time, trivial contacts due to short sequence separations \( \Delta < 100 \text{ bp} \) are not counted. We note that at this stage, and at the next one too, there could be overlaps in the regions covered by different putative plectonemes.

Within each band with \( \Delta < 400 \text{ bp} \), we then searched for the presence of tight contacts, a signature feature of superhelices. Specifically, we searched for the contacts at distance smaller than 7.5\( \sigma \) (\( \sim 6 \text{ Å} \)), and took the contact map diagonal. These bands correspond to dsDNA stretches in spatial proximity and with opposite directionality (in an oriented ring).

For each band we identified its apex, which is the site, \( i \), associated to the longest uninterrupted ladder of contacts \( \{i + \Delta, i - \Delta\}, (i + \Delta + 1, i - \Delta - 1), (i + \Delta + 2, i - \Delta - 2) \), starting at a sequence separation, \( \Delta \), not larger than 400 \text{ bp} to avoid excessively large apical loops. At the same time, trivial contacts due to short sequence separations \( \Delta < 100 \text{ bp} \) are not counted. We note that at this stage, and at the next one too, there could be overlaps in the regions covered by different putative plectonemes.

Within each band with \( \Delta < 400 \text{ bp} \), we then searched for the presence of tight contacts, a signature feature of superhelices. Specifically, we searched for the contacts at distance smaller than 7.5\( \sigma \) (\( \sim 6 \text{ Å} \)), and took the contact map diagonal. These bands correspond to dsDNA stretches in spatial proximity and with opposite directionality (in an oriented ring).

For each band we identified its apex, which is the site, \( i \), associated to the longest uninterrupted ladder of contacts \( \{i + \Delta, i - \Delta\}, (i + \Delta + 1, i - \Delta - 1), (i + \Delta + 2, i - \Delta - 2) \), starting at a sequence separation, \( \Delta \), not larger than 400 \text{ bp} to avoid excessively large apical loops. At the same time, trivial contacts due to short sequence separations \( \Delta < 100 \text{ bp} \) are not counted. We note that at this stage, and at the next one too, there could be overlaps in the regions covered by different putative plectonemes.

Within each band with \( \Delta < 400 \text{ bp} \), we then searched for the presence of tight contacts, a signature feature of superhelices. Specifically, we searched for the contacts at distance smaller than 7.5\( \sigma \) (\( \sim 6 \text{ Å} \)), and took the contact map diagonal. These bands correspond to dsDNA stretches in spatial proximity and with opposite directionality (in an oriented ring).
coiled configurations except for a small subset (0.5% of 51-knotted instances and even smaller for 52-knotted and unknotted ones) with non well formed superhelical regions or atypically large apical loops, see Supplementary Figure S3.

**Topological observables**

To locate the knotted region along the ring we used the bottom-up search scheme described in ref. (58). The procedure consists of searching for the smallest portion of the ring (starting from portions of only few nucleotides and systematically expanding to longer ones) that, after closure, has the same topology of the entire ring. The topology was established using Alexander polynomials evaluated at $t = -1$ and $t = -2$. For closing the considered portion we used the minimally interfering closure scheme (58), where the termini of the portion are bridged either with a straight segment or via a path involving the convex hull, depending on their proximity.

**RESULTS**

**Conformational variability**

We first discuss the conformational variability observed in simulations of 2 kbp-long knotted DNA rings with the 5% negative supercoiling typical of bacterial plasmids. Respect to earlier studies on 31-knotted DNAs, we stepped up the complexity and considered the 51 and 52 topologies, respectively a torus and a twist knot. With these more complex knots we can explore the effects of a larger writhe and more numerous minimal crossings on the branchedness and dynamics of DNA rings. In addition, 5-crossing knots provide the simplest context for an equal footing comparison of torus and twist knots, which typically show different physical behavior, from mechanical resistance to sliding friction to pore-translocation compliance (59,60).

Typical snapshots, representative of the various degrees of branching found in the 51- and 52-knotted DNA rings, are given in Figure 2A along with instances without knots (01 case). The observed conformational variability is significant and is consistent with that recently reported for shorter unknotted plasmids based on cryo-em experiments and atomistic simulations (29).

It is interesting to examine the relationship between knottedness and the number of branches, or plectonemically-wound regions, because of the competing elements that govern it. The lobes, apices and clasps inherent to complex knots can favour plectonemes by serving as nucleation points, while the conformational restrictions of the topological constraints can inhibit plectonemes’ formation. The histograms in Figure 2B clarify that, at this contour length, unknotted DNA rings are actually somewhat richer in plectonemes than knotted ones; in particular, instances with three or more plectonemes are practically found in unknotted rings only.

More conspicuous differences related to topology are found in the distributions of plectoneme lengths, $l_{ple}$, and gyration radius, $R_g$, see Figure 3A and B. In particular, the conditional distributions of $l_{ple}$ for the common single- and double-plectoneme states are little superposed for unknotted rings, but overlap substantially for knotted ones, see Supplementary Figure S4 for the 52 topology. The length of the plectonemes is also different across the 2kbp-long knotted and unknotted rings. For examples, plectonemes longer than 1500 bp are common in unknotted rings but rare in 51-knotted ones (50.7% and 0.05% of the populations, respectively). Conversely, conformers with only one plectoneme, and shorter than 1000 bp, are uncommon in unknotted rings but abundant in 51-knotted ones (6.3% and 22% of the populations, respectively).

These differences could be of practical relevance, since they could be exploited in imaging experiments to tell apart knotted from unknotted plasmids when supercoiling is present. Such discrimination is generally beyond the scope of gel electrophoresis, the method of choice for DNA topological profiling, but for a notable exception see (61).

The interplay of knot length, $l_k$, plectoneme length and gyration radius is presented in Figure 3C and D. The joint
probability distribution in panel C shows that $l_k$ and $R_g$ are anticorrelated, see also Supplementary Figure S5. What is specific of supercoiled knotted DNA rings is, instead, the presence of two peaks in the joint $l_k - R_g$ distribution. The peaks’ origin is clarified by their marginal (projected) $l_k$ distributions subdivided by number of plectonemes, see Figure 3D. Specifically, the dominant peak, for $l_k \sim 700$ bp, is mostly associated to single-plectoneme states, while the peak at larger knot lengths ($l_k \sim 1100$ bp), corresponds to states with two or more plectonemes. Analogous results for the $S_2$ topology are shown in Supplementary Figure S6.

The inverse correlation of $l_k$ and the number of plectonemes is understood by noting that the knotted region, which is the shortest uninterrupted portion of the ring accommodating the essential crossings (the knotted core), must also include all intervening loops between the crossings except for the longest one. In supercoiled rings this remainder loop typically coincides with a plectoneme. Because the average plectoneme length decreases as they become more numerous, one has that $l_k$ is shorter for states with a single plectoneme.

**Time evolution of metric and knot-related properties**

The data shown in Figure 4A are a kinetic counterpart to the static, or ensemble, view given above of the interplay of the knotted region, the number of plectonemes and the gyration radius.

One notes that over the typical duration of a trajectory ($2.2 \times 10^6 \tau_{LJ}$ corresponding to $\sim 1.5$ ms) both $l_k$ and $R_g$ have significant fluctuations, and clearly of opposite sign. These are accompanied by several changes in the number of plectonemes, as conveyed by the coloured background.

A more quantitative analysis of the characteristic timescales of these variations is given in panels B and C of Figure 4. These panels present the autocorrelation curves of $R_g$, $l_k$ and of $l_{plc}$. The latter, was used in place of the number of plectonemes (on which it clearly depends) for its broader range of values, which makes it more amenable to the autocorrelation analysis. By integrating the autocorrelation curves, one has that the characteristic times of $l_k$ and $R_g$ are respectively equal to $1.03 \times 10^6 \tau_{LJ}$ and $0.53 \times 10^6 \tau_{LJ}$, while for plectonemes’ length it is $0.69 \times 10^6 \tau_{LJ}$. Consistent with visual inspection, these timescales are all of the same order, about $10^6 \tau_{LJ}$, which is much shorter than the duration of each simulated trajectory.

It is interesting that unknotted rings have a somewhat slower internal kinetics than knotted rings, cf. panels B and C in Figure 4. In fact, the characteristic times of $R_g$ and $l_{plc}$ are, respectively 80% and 35% longer for unknotted rings (the same holds for $R_g$ in the torsionally-relaxed case, see Supplementary Figure S7). The result is not obvious, as one might expect a slower internal dynamics for knotted rings due to the friction of their self-contacts. It can be explained by considering that a finite portion of topological constraints necessarily uses up a finite portion of the chain, and therefore knotted rings have a shorter effective contour length than unknotted ones and a smaller gyration radius too (see Figure 3). This, in turn, reflects in a reduced breadth of the reveal conformational space and hence a faster relaxation kinetics.

From the above analysis of overall metric and topological properties we conclude that, at physiological supercoiling, 2 kbp-long knotted rings have enough conformational freedom to fluctuate spontaneously between two main states, related to the peaks in Figure 3C, differing by knot size as well as the length and number of plectonemes. The characteristic timescales of these variations is $\sim 3 \times 10^6 \tau_{LJ}$, corresponding to about 0.02 ms, see Figure 4A and Supplementary Figure S8.

**Slowly-moving boundaries of the knotted region**

To better understand the kinetics of the concerted variations of knot size and the number of plectonemes, we examined the time evolution of knotted and supercoiled regions, as in the kymographs of Figure 5A, for other examples see Supplementary Figure S9.

The kymograph clearly indicates that there exists an additional relevant kinetic process besides those discussed before, namely a surprisingly slow stochastic motion of the knot along the ring contour.
Knotting and supercoiling are both required for persistent interlockings

To address this point, we decoupled entanglement and supercoiling by studying the time evolution of DNA rings where either the knot or supercoiling were present, but not both of them.

The results are shown in the kymographs of panels B and C of Figure 5. Their properties are in stark contrast with those of panel A, where knotting and supercoiling were both present. In fact one observes that neither the boundaries of the knotted region (without supercoiling) nor those of plectonemes (without knotting) are inherently persistent. In fact, in both cases the boundaries vary on the same relatively fast timescales of the metric relaxation, i.e. $\sim 10^6\tau_{LJ}$.

We thus conclude that it is precisely the synergistic action of complex topology and supercoiling that is responsible for the locked boundaries, and the latter disappear when either of them is missing.

To clarify the mechanism underpinning this effect we inspected in detail the dynamical evolution of the rings. We thus established that the persistent boundaries correspond to specific points of tight and complex self-contacts of the knotted region.

A large number of self-contacting loci are clearly introduced by supercoiling in any DNA ring, regardless of its topological state. In knotted rings, due to the tightening of the intrinsic essential crossings, these loci typically involve several clasped or hooked double strands, as highlighted in the snapshots of Figure 5A. We observed that, similarly to what happens during the pore translocation of knotted filaments (59,60,64,65), the topological friction at these points is so high that the DNA strands are locally pinned while other parts of the chain can reconfigure.

This effect accounts for the observed separation of timescales between the metric relaxation time and the contour motion of the clasped points, at the boundaries of the knotted region. Incidentally, we note that the relevance of these persistent regions of self-contacts reinforces a posteriori the necessity to use models, such as oxDNA, where the spatial description is sufficiently fine to capture the internal friction that develops when two tightly interacting DNA strands slide against each other. For this reason, we surmise that the observed sliding hindrance of contacting

Note, in fact, that the region covered by the knot at the beginning of the simulation in Figure 5A (bp 600–bp 1800) still has not moved appreciably by the end of the trajectory (bp 700–bp 2000). Considering that the trajectory covers a timespan of $2 \times 10^7\tau_{LJ} \sim 1.5$ ms, it is clear that the contour motion of the knotted region occurs over timescales that are much longer than those discussed previously. These, we recall, were the relaxation times of the gyration radius and knot length and the switching time between tight and de-localised knotted states. The former two were about equal to $10^5\tau_{LJ}$ while the latter was about $3 \times 10^5\tau_{LJ}$, see Figure 4 and Supplementary Figures S7 and S9.

We thus conclude that in supercoiled DNA rings, the contour motion of the knotted region is slower than these other processes by an order of magnitude (and possibly more since our simulations had a typical maximum duration of $\sim 2 \times 10^6\tau_{LJ}$).

Besides the knot, Figure 5A shows the concurrent evolution of plectonemically-wound regions, too. Note that most of the time, there is a single long plectoneme that spans the ring portion complementary to the knotted region; but the latter can occasionally nest a plectoneme too. The kymograph clarifies that such nested instances have the following properties: (i) they occur in addition, and not in substitution, of the typically longer ‘dominant’ plectoneme following properties: (i) they occur in addition, and not in substitution, of the typically longer ‘dominant’ plectoneme that complements the knotted region; (ii) their characteristic lifetime is $7 \times 10^5\tau_{LJ}$ and (iii) consecutive appearances are separated by intervals of highly variable duration.

Overall, the several observed fluctuations of plectonemes’ number and length during the entire trajectory are in line with the relatively fast relaxation dynamics of $R_g$ and $l_{plc}$ of Figure 4. Strikingly, these variations are accompanied by a noticeable persistence of the plectoneme boundaries, which mirrors the one of the knotted region.

These features are ubiquitous across the collected trajectories for both $5_1$ and $5_2$ topologies, see Supplementary Figure S9. One concludes that, irrespective of their torus ($5_1$) or twist ($5_2$) character, these supercoiled knotted rings have persistent boundaries between knotted and plectonemically-wound regions. This, in turn, poses the question of which of these two components, knots or plectonemes, is the primary cause for the slow evolution of these boundaries.

Knotting and supercoiling are both required for persistent interlockings

To address this point, we decoupled entanglement and supercoiling by studying the time evolution of DNA rings where either the knot or supercoiling were present, but not both of them.

The results are shown in the kymographs of panels B and C of Figure 5. Their properties are in stark contrast with those of panel A, where knotting and supercoiling were both present. In fact one observes that neither the boundaries of the knotted region (without supercoiling) nor those of plectonemes (without knotting) are inherently persistent. In fact, in both cases the boundaries vary on the same relatively fast timescales of the metric relaxation, i.e. $\sim 10^6\tau_{LJ}$.

We thus conclude that it is precisely the synergistic action of complex topology and supercoiling that is responsible for the locked boundaries, and the latter disappear when either of them is missing.

To clarify the mechanism underpinning this effect we inspected in detail the dynamical evolution of the rings. We thus established that the persistent boundaries correspond to specific points of tight and complex self-contacts of the knotted region.

A large number of self-contacting loci are clearly introduced by supercoiling in any DNA ring, regardless of its topological state. In knotted rings, due to the tightening of the intrinsic essential crossings, these loci typically involve several clasped or hooked double strands, as highlighted in the snapshots of Figure 5A. We observed that, similarly to what happens during the pore translocation of knotted filaments (59,60,64,65), the topological friction at these points is so high that the DNA strands are locally pinned while other parts of the chain can reconfigure.

This effect accounts for the observed separation of timescales between the metric relaxation time and the contour motion of the clasped points, at the boundaries of the knotted region. Incidentally, we note that the relevance of these persistent regions of self-contacts reinforces a posteriori the necessity to use models, such as oxDNA, where the spatial description is sufficiently fine to capture the internal friction that develops when two tightly interacting DNA strands slide against each other. For this reason, we surmise that the observed sliding hindrance of contacting

Note, in fact, that the region covered by the knot at the beginning of the simulation in Figure 5A (bp 600–bp 1800) still has not moved appreciably by the end of the trajectory (bp 700–bp 2000). Considering that the trajectory covers a timespan of $2 \times 10^7\tau_{LJ} \sim 1.5$ ms, it is clear that the contour motion of the knotted region occurs over timescales that are much longer than those discussed previously. These, we recall, were the relaxation times of the gyration radius and knot length and the switching time between tight and de-localised knotted states. The former two were about equal to $10^5\tau_{LJ}$ while the latter was about $3 \times 10^5\tau_{LJ}$, see Figure 4 and Supplementary Figures S7 and S9.

We thus conclude that in supercoiled DNA rings, the contour motion of the knotted region is slower than these other processes by an order of magnitude (and possibly more since our simulations had a typical maximum duration of $\sim 2 \times 10^6\tau_{LJ}$).

Besides the knot, Figure 5A shows the concurrent evolution of plectonemically-wound regions, too. Note that most of the time, there is a single long plectoneme that spans the ring portion complementary to the knotted region; but the latter can occasionally nest a plectoneme too. The kymograph clarifies that such nested instances have the following properties: (i) they occur in addition, and not in substitution, of the typically longer ‘dominant’ plectoneme that complements the knotted region; (ii) their characteristic lifetime is $7 \times 10^5\tau_{LJ}$ and (iii) consecutive appearances are separated by intervals of highly variable duration.

Overall, the several observed fluctuations of plectonemes’ number and length during the entire trajectory are in line with the relatively fast relaxation dynamics of $R_g$ and $l_{plc}$ of Figure 4. Strikingly, these variations are accompanied by a noticeable persistence of the plectoneme boundaries, which mirrors the one of the knotted region.

These features are ubiquitous across the collected trajectories for both $5_1$ and $5_2$ topologies, see Supplementary Figure S9. One concludes that, irrespective of their torus ($5_1$) or twist ($5_2$) character, these supercoiled knotted rings have persistent boundaries between knotted and plectonemically-wound regions. This, in turn, poses the question of which of these two components, knots or plectonemes, is the primary cause for the slow evolution of these boundaries.
strands would be even higher in atomistic or more fine-grained models at this same high salt conditions. At the same time, the friction between DNA strands could be relieved by increasing their electrostatic repulsion with a lower salt concentration. These would be worthwhile points to address in future modelling studies and possibly also experimentally, e.g. using setups akin to those of ref. (66).

Importantly, not all intrinsic (or essential) crossings of the knots create the persistent interlocking, but only a subset of them. This is visible in the snapshots of Figure 5A where the two boundaries of the five-crossing knot are pinned by their high local physical entanglement and yet the DNA strands can slide internally of the knotted region, despite the several points of pairwise contacts. It is precisely this internal sliding that creates the possibility for plectonemes to form transiently, but repeatedly, within the knotted region. As a matter of fact, the persistent interlocking appears in either of the two qualitatively different conformers populated by supercoiled knotted rings (i.e. those with local or non-local knots in Figure 3).

In this regard, it is relevant to recall the seminal work of Liu et al. (38,39), who pointed out that hooked DNA juxtapositions are an ideal target substrate of topoII enzymes because local strand passages at these point generally produces a simpler topology. The kinetic persistence of these multi-strand interlockings adds a novel temporal dimension to other, more thermodynamical effects of supercoiling, such as knot localization, that are credited to favour the local, yet globally-disentangling action, of topo II. The present results, in fact, complement the insight from earlier thermodynamics sampling (43,44), by showing that once hooked or multiply-clasped juxtapositions are formed, they are long-lived. This, we speculate, is key for making such forms of local entanglement persistent enough to be recognised and processed by topo II enzymes.

**DISCUSSION**

In summary, we used molecular dynamics simulation and the oxDNA mesoscopic model to study the effect of complex, five-crossing knots, on the conformational and kinetic properties of 2kbp-long plasmids with the typical 5% negative supercoiling found in bacterial plasmids. We particularly focussed on whether and how complex topologies, with their numerous points of high curvature and self-contact,
can alter the branchedness of supercoiled plasmids and the dynamical evolution at long timescales.

On both accounts, we found that the interplay of knotting and supercoiling has major consequences that would have been difficult to anticipate apriori.

For the structural properties, we found that the conformational ensemble explored during the spontaneous dynamical evolution is largely dominated by two qualitatively-distinct states. They differ both by knot size and degree of branching. This fact, noteworthy per se, is accompanied by two intriguing kinetic effects. The first is that spontaneous fluctuations between these two states occur on timescales that are comparable to metric relaxation times of unknotted rings. The second is that certain boundaries separating the knotted and plectonemically-wound regions are very long-lived, and remain persistent over timescales that are much longer, arguably by at least an order of magnitude, than the metric relaxation times.

This complex phenomenology is shown to arise exclusively from the cooperative action of supercoiling and topological constraints; removing either of the two suffices to remove the persistent boundaries. The latter are shown to occur in correspondence of loci where multiple strands become interlocked. The interlockings have a local geometry that is analogous to the so-called hooked juxtapositions (38–40), argued to be ideal local targets for the topoisomerases’ knot simplifying action. We accordingly surmise that their long-lived character, besides their structural features, could also be instrumental for their recognition by topoisomerases. We believe this would be a noteworthy problem to address in future studies, for instance using mesoscale models incorporating the interaction of DNA and proteins, which would be essential for a realistic description of DNA organization and processing in vivo. In addition, we expect that long-lived multi-strand interlockings could be probed with single-molecule manipulation techniques such as pore translocation, which has been previously used on DNA rings with either knots or supercoiling (48,65,67–71).

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

**FUNDING**

Italian Ministry of Education. Funding for open access charge: Institutional funds. Conflict of interest statement. None declared.

**REFERENCES**


