How topoisomerase IV can efficiently unknot and decatenate negatively supercoiled DNA molecules without causing their torsional relaxation

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ABSTRACT

Freshly replicated DNA molecules initially form multiply interlinked right-handed catenanes. In bacteria, these catenated molecules become supercoiled by DNA gyrase before they undergo a complete decatenation by topoisomerase IV (Topo IV). Topo IV is also involved in the unknotting of supercoiled DNA molecules. Using Metropolis Monte Carlo simulations, we investigate the shapes of supercoiled DNA molecules that are either knotted or catenated. We are especially interested in understanding how Topo IV can unknot right-handed knots and decatenate right-handed catenanes without acting on right-handed plectonemes in negatively supercoiled DNA molecules. To this end, we investigate how the topological consequences of intersegmental passages depend on the geometry of the DNA-DNA juxtapositions at which these passages occur. We observe that there are interesting differences between the geometries of DNA-DNA juxtapositions in the interwound portions and in the knotted or catenated portions of the studied molecules. In particular, in negatively supercoiled, multiply interlinked, right-handed catenanes, we detect specific regions where DNA segments belonging to two freshly replicated sister DNA molecules form left-handed crossings. We propose that, due to its geometrical preference to act on left-handed crossings, Topo IV can specifically unknot supercoiled DNA, as well as decatenate postreplicative catenanes, without causing their torsional relaxation.

INTRODUCTION

Bacterial DNA topoisomerase IV (Topo IV) is responsible for the decatenation of freshly replicated circular DNA molecules (1,2). Topo IV belongs to the type II DNA topoisomerases, which perform passages of two duplex DNA segments through each other. During the reaction, Topo IV binds first to one DNA region and then drives the passage of another DNA region through a transient cut introduced in the first region. After the passage, the transient cut is resealed (3). The passage is driven by the energy gained from ATP hydrolysis and is orchestrated by a complex sequence of conformational changes in the enzyme. Topo IV has to solve a particularly difficult problem. On one hand, Topo IV has to perform intersegmental passages resulting in the decatenation of freshly replicated circular double-stranded DNA molecules that wind around each other in a right-handed way (4–9). On the other hand, Topo IV should not perform intersegmental passages within interwound portions of supercoiled DNA molecules where the opposing DNA segments also wind around each other in a right-handed way (5). Locally, the structure of the regions where the catenated DNA molecules wind around each other are very similar to the plectonemic regions in negatively supercoiled DNA (see Figure 1). However, the topological consequences of passages are different. Passages between duplex segments belonging to two different molecules lead to their progressive decatenation, which is necessary for segregation of freshly replicated DNA molecules. On the other hand, intramolecular passages between duplex segments in plectonemically wound negatively supercoiled DNA molecules...
lead to their progressive relaxation, which is an undesired effect. Supercoiling is actively maintained by bacterial cells since, among other roles it plays, it is required to facilitate DNA replication and transcription (10). If Topo IV were efficient in relaxing negatively supercoiled DNA molecules, this relaxation would result in a metabolic short circuit. The relaxation of negatively supercoiled DNA molecules would be followed by the action of DNA gyrase that, using the energy gained from ATP hydrolysis, would act to reestablish the original level of negative supercoiling (11). Negatively supercoiled molecules then again would be relaxed by topoisomerase IV in a reaction that also uses ATP. Such a metabolic short circuit would be harmful for cells as it would deplete the available ATP.

It has not yet been firmly established how Topo IV distinguishes between right-handed intertwinings of two duplex DNA regions in catenanes versus intertwinings in supercoiled DNA molecules.

In addition to its role in the decatenation of postreplicative catenanes, Topo IV (1,2) also plays an important role in the relaxation of positive supercoils arising ahead of active replication forks and also ahead of transcribing RNA polymerases (12,13). Without efficient relaxation of positive supercoiling, DNA replication and transcription would be strongly inhibited (12,14). Experiments performed in vitro showed that Topo IV is ~20 times more active in the relaxation of positive supercoiling than in the relaxation of negative supercoiling (15). It has been proposed that these differences in the activity on positively and negatively supercoiled DNA result from the enzyme’s preference for DNA juxtapositions in which the juxtaposed segments wind around each other in a left-handed way and have the geometry typical for the plectonemes formed in positively supercoiled DNA (4). In principle, such a preference should make Topo IV inefficient in resolving right-handed postreplicative catenanes. However, this is not the case. The puzzle as to how Topo IV can act efficiently on left-handed duplex-duplex windings in positively supercoiled DNA and still can be active in resolving right-handed postreplicative catenanes is known as the Topo IV paradox (6).

Topo IV also has the ability to unknot DNA molecules preferentially and thus to decrease the knotting level much below the level that would result from random intramolecular passages (16). While unknotting of DNA molecules may seem less vital than the decatenation of freshly replicated DNA molecules, DNA knots need to be removed efficiently by DNA topoisomerases or else the knots would interfere with transcription and DNA replication (17,18). In fact, DNA knots frequently form during the replication of circular DNA (19). Unknotting and decatenation of freshly replicated DNA molecules occur concomitantly in vivo and Topo IV is the enzyme involved in the two processes (20). The preferential targeting of DNA knots by Topo IV was proposed to result from enzyme selectivity for the juxtapositions involving strongly bent DNA (21–24). This proposal was confirmed when the solved structure of Topo IV bound to DNA revealed that the bound G segment is strongly bent within the protein (3). The G segment is the DNA region that is first bound by a topoisomerase and then cut to serve as the Gate for the passage of the T segment that is Transferred through the gate.

In bacterial cells, the DNA is bent significantly due to topological constraints such as supercoiling or catenation (see Figure 1). When Topo IV, binds to DNA, which is already bent, the lowest energy state is reached when the enzyme positions itself in such a way that the DNA bending due to supercoiling, knotting, or catenation is best accommodated into the enzyme binding site, which imposes DNA bending (25). The azimuthal orientation of Topo IV on the bent G segment is such that it only can capture potential T segments that are inside of the bend formed by the G segment (3). For the DNA-DNA passage, the T segment needs to be captured by the Topo IV before passing through the entry and exit gates of the enzyme. The passages can happen efficiently when the G segment, which determines the spatial orientation of Topo IV, and the T segment are in a suitable angular inclination with respect to each other. The dependence of Topo IV activity on the inclination or crossing angle (θ) between the G and T segment was investigated earlier (4,6,15). However, these earlier studies did not consider cases where the torsional tension due to supercoiling was acting on knotted and catenated DNA molecules as is the case of DNA in bacterial cells.

Topo IV’s geometrical/structural selectivity for intersegmental passages was presumably under strong evolutionary pressure until it reached the best compromise between three requirements: (i) selective decatenation of negatively supercoiled DNA molecules without causing their torsional relaxation; (ii) selective unknotting of supercoiled DNA molecules without causing their torsional relaxation; (iii) ef-

![Figure 1. Topo IV needs to distinguish between right-handed intertwinings of DNA duplexes resulting from postreplicative catenation and those resulting from negative supercoiling. A schematic presentation of supercoiled postreplicative catenanes is shown in the center. Two newly replicated circular DNA molecules (red and blue) form multiply interlinked right-handed catenanes. The molecules also are supercoiled, which results in the formation of right-handed plectonemes. Local structural cues guide Topo IV to perform passages resulting in progressive decatenation (green arrow) and to avoid passages within the interwound portions (red arrows) that would result in a harmful relaxation of the negative supercoiling.](https://academic.oup.com/nar/article-abstract/44/10/4528/2516939)
sufficient relaxation of positively supercoiled DNA but not of negatively supercoiled DNA.

Using numerical simulations, we investigate the possible geometrical selectivity of Topo IV that satisfies these three requirements.

MATERIALS AND METHODS

Metropolis Monte Carlo

We model circular DNA chains of length 3 kb and diameter 3 nm with bending and torsional stiffness. The diameter of 3 nm represents the effective diameter of DNA under conditions where DNA charges are strongly, but not completely, screened, such as is the case \textit{in vivo} (26–28). Supercoiling was introduced by imposing various levels of linking number difference ($\Delta Lk$).

DNA chains were modeled as circular beaded chains, each with $N$ hard-core beads of diameter $\sigma$ centered at the vertices $r_i$ of an equilateral polygon having $N$ edges of length $\sigma$. For a given chain configuration, the total energy is defined as

$$E = E_V + E_B + E_T.$$ 

The hard-core steric repulsion is given by

$$E_V = \begin{cases} 0 & \text{if } |r_i - r_j| > \sigma \text{ for all nonconsecutive } i, j \\ \infty & \text{otherwise} \end{cases}.$$ 

The bending stiffness is imposed by the usual bending energy term

$$E_B = \frac{k}{2} \sum_i \beta_i^2,$$

where $\beta_i$ is the bending angle

$$\cos(\beta_i) = \frac{(r_{i+1} - r_i) \cdot (r_i - r_i-1)}{|r_{i+1} - r_i| |r_i - r_i-1|}.$$ 

Torsional stiffness is imposed by the energy term proposed by Vologodskii \textit{et al.} (29)

$$E_T = \frac{2\pi^2}{L} C(\Delta Lk - Wd)^2,$$

with $L = N\sigma$ the chain length, $\Delta Lk$ the delta linking number, and $Wd$ the writhe of the chain (30).

To model 3 kb DNA chains, assuming a linear density of 3 bp/nm, we used the parameters listed below:

- $N = 334$
- $\sigma = 3$ nm
- $k = 17k_B T$
- $C = 72k_B T$ nm
- $\Delta Lk = -16$.

The bending stiffness parameter $k$ was chosen to give a persistence length $L_p \approx 53$ nm for a linear chain without torsional stiffness. The torsional rigidity constant $C$ was taken in the range proposed by Vologodskii \textit{et al.} (29). The linking number difference $\Delta Lk$ was set to $-16$ since this value places simulated molecules in the middle of the supercoiling density ranges observed in DNA plasmids isolated from bacteria grown in various conditions (31). As mentioned in the discussion, the magnitude of supercoiling is reduced \textit{in vivo}, but this change is balanced by the crowding effect (32).

To evaluate thermal equilibrium properties of the system, we generated an ensemble of chain configurations, distributed according to the Boltzmann distribution, using a Metropolis Monte Carlo algorithm (33) based on crankshaft moves, described earlier (34). Note that this type of crankshaft move conserves the distance between consecutive beads and the topology of the chain by preventing the rotating sub-chain from crossing other parts of the chain during the move.

For the knotted chains, after an initialization starting from the configuration given in Supplementary Figure S1A, the Monte Carlo algorithm was run for a total of $2.6 \times 10^8$ iterations (one iteration = $N$ crankshaft moves, where $N$ is the number of beads). After the first $10^7$ iterations, sample configurations were saved every $10^5$ iterations to be used later to evaluate the thermal averages of properties of interest. During the run, the squared radius of gyration of knotted supercoiled chains was measured every 1000 iterations, and the corresponding autocorrelation function was evaluated using the ‘blocking’ method (35). Supplementary Figure S2 (red points) presents the autocorrelation as a function of block size. A fit of exp$(-n/\tau)$ to these data gives an estimated correlation time $\tau$ for the squared radius of gyration on the order of $10^6$ iterations, which means that the total number of Monte Carlo iterations corresponded approximately to 2600$\tau$. This result suggests that enough independent configurations were obtained during the Monte Carlo run to sample thermal equilibrium.

For the two catenated chains, the system was initialized to the configuration shown in Supplementary Figure S1B, and the Monte Carlo algorithm was run for a total of $3.5 \times 10^8$ iterations. Sample configurations were saved every $10^5$ iterations after the first $10^7$ iterations. The autocorrelation function for the average squared radius of gyration per chain is shown in Supplementary Figure S2 (blue points), and the estimated correlation time is on the order of $2 \times 10^5$ iterations. Therefore, the total number of iterations corresponded approximately to 1700$\tau$, which suggests a sufficiently equilibrated system of catenated, supercoiled chains.

Geometrical and topological characterization of DNA-DNA juxtapositions

In our simulations, DNA molecules are modeled as beaded chains where every bead corresponds to 8 bp. To analyze the juxtapositions, in which the G and T segments have sizes of about 50 bp, we analyzed subchains composed of seven beads each. For the geometrical analysis each seven bead-long subchain of modeled configurations was considered as a potential G segment and its geometry was approximated by a fixed angle bend formed by the two line segments starting at the center of the middle bead (M1) and ending at the two centers of the terminal beads (see Figure 2). As explained earlier, Topo IV positions itself on a G segment in such a way that it can only capture potential T segments that are located inside of the bend formed by the G segment. To account for this type of Topo IV’s selectivity, we entered into the statistics only the DNA–DNA juxtapositions where the subchain S2 (representing the potential T segment) had an
intersection point \( I_p \) with the surface spanning from the interior of the fixed angle bend that approximated the geometry of the G segment (see Figure 2). In addition, to reflect the fact that topoisomerases can only capture DNA segments that are very close to its surface, we somewhat arbitrarily decided that the distance between the middle bead \( M_1 \) of the G segment and the intersection point \( I_p \) should not exceed 6 nm (i.e. \( 2 \sigma \)). If there was more than one intersection point with the spanning surface determined by the bend of a given G segment, only the one closest to the vertex \( M_1 \) was taken into account. In such a case the seven bead-long subchain \( S_2 \) representing a potential T segment is then defined as the one whose middle vertex \( M_2 \) is the vertex closest to the \( I_p \). Subsequently, for the geometrical analysis, the subchain \( S_2 \) was replaced by the fixed angle bend formed by the two line segments starting from vertex \( M_2 \) and ending at the two terminal vertices of that chain. Once the pairs of potential G and T segments were defined, we characterized them geometrically and considered how the topology of the entire analyzed molecules changes upon intersegmental passage between these G and T segments.

**Geometrical analysis.** We characterized several geometric characteristics of the analyzed juxtapositions. However, the ones that are likely to be used by topoisomerases in the selection of juxtapositions for strand passage, and were considered as such in several earlier papers (4,6,21,24), are the crossing angle \( \theta \) and the bending angle \( \alpha_1 \). The crossing angle \( \theta \) measures the inclination of the G segment, which is bound first by the topoisomerase and whose curvature, reflected by the angle \( \alpha_1 \), dictates how the topoisomerase is positioned on the DNA. This positioning, in turn, determines from which side and at which crossing angle \( \theta \), the T segment (subchain \( S_2 \)) can approach the G segment to be used for the passage. The geometries of seven bead-long subchains are approximated by fixed angle bends formed by two line segments connecting the center of the middle beads with the centers of the terminal beads. For each juxtaposition (see the Materials and Methods section for our criteria of juxtapositions), we determine the \( \alpha_1 \) angle and the chirality-dependent crossing angle \( \theta \). In the inset the tangentials \( N_2 \) and \( N_1 \) are shown along the direction that is perpendicular to both of them. In that projection, the \( \alpha_1 \) angle and the angle of the tangential closer to the observer (in this case \( N_2 \)). If the direction of this rotation is clockwise, the crossing is considered as right-handed and its angular value is given a positive sign. The \( \theta \) angle can range from \(-90^\circ \) to \(+90^\circ \). Where negative and positive values indicate left- and right-handed crossings, respectively, and where the \( 0^\circ \) value characterizes achiral, perpendicular crossings. It is important to mention that the \( \alpha_1 \) angle notation used here is different than the \( \alpha_1 \) angle used by Stone et al. (4) and results in a \(-90^\circ \) shift. In our notation, right-handed and left-handed crossings have positive and negative \( \alpha_1 \) values, respectively. The convention used by Stone et al. associates a positive sign to all \( \theta \) angles and they range from \( 0^\circ \) to \( 90^\circ \) for left-handed crossings and from \( 90^\circ \) to \( 180^\circ \) for right-handed crossings.

The \( \alpha_1 \) bending angle, reflecting the bending angle of the G segment bound by a topoisomerase, is simply the included angle (\( \leq 180^\circ \)) of a given bend that approximates the geometry of a given seven bead-long subchain (see Figure 2).

**Topological analysis.** For each geometrically characterized juxtaposition, we determined how intersegmental passage within the given juxtapositions changes the topology of the entire modeled DNA molecule. To this end, for every analyzed juxtaposition, we created a configuration resulting from a passage between juxtaposed segments. In the case of knotted, supercoiled molecules, we calculated the HOMFLY polynomial to determine the knot type of the configuration after the intersegmental passage (36,37). For supercoiled, catenated molecules, we calculated the catenation number, i.e. the linking number between the two molecules forming the catenane, to determine the topology of the catenane after the intersegmental passage.

**RESULTS**

**Structure of negatively supercoiled DNA molecules forming right-handed trefoil knots**

Trefoil knots are the simplest and the most frequent knots that can be formed in DNA plasmids in vitro (26,38) and also in vivo (19,39). Trefoil knots exist in right- and left-handed forms. In right-handed trefoil knots (+\( z_1 \) in standard knot notation), the DNA–DNA juxtapositions due
Visual analysis of DNA molecules shown in Figure 3 supports the idea that portions of knotted supercoiled DNA molecules are moving between the two apical loops. Crossings between approaching segments in the supercoiled DNA molecules (Figure 3A) are right-handed with the knotted portion moving between the two apical loops.

Crossing angle differences between knotted and interwound portions of supercoiled DNA molecules

Visualization of DNA molecules shown in Figure 3 suggests that crossings between approaching segments in the knotted portions are closer to perpendicular than the crossings in the plectonemic portion of the molecules. To verify this observation, we measured the crossing angle $\theta$ for DNA–DNA juxtapositions, where the potential T segment was inside the bend formed by the potential G segment and determined the topological consequences of passages occurring at these juxtapositions (see Figure 2 and the Materials and Methods section for more details).

Figure 4A shows the distribution of the chirality-dependent $\theta$ angles in DNA-DNA juxtapositions observed in simulated negatively supercoiled DNA molecules forming right-handed trefoil knots, examples of which are shown in Figure 3. The height of the main histogram bars in Figure 4A corresponds to the observed number of juxtapositions within the corresponding $\theta$ angle range in the simulated DNA molecules. We can see that the most frequent juxtapositions are right-handed with the $\theta$ angle ranging between 50° and 60°. Such juxtapositions are expected for DNA molecules where the duplex regions wind around each other in a right-handed fashion (4). As can be seen in Figure 3, most of the DNA–DNA juxtapositions in these molecules are between duplexes forming the interwound portions of the molecules. The passages occurring between the interwound duplexes result in the undesired partial relaxation of DNA supercoiling and have no effect on knotting or unknotting of these molecules. The fraction of juxtapositions with this topological outcome is indicated in green within the individual histogram bars. The blue color indicates the fraction of juxtapositions that, upon internal passage, result in DNA unknotting. These juxtapositions can only form in the knotted portion of the molecules.

The topological outcome of passages occurring in given juxtapositions is therefore a strong hint or proxy for the localization of these juxtapositions and we use it as such in what follows. The main histogram in Figure 4A reveals that there is a difference in the distribution of the $\theta$ angle in the interwound and knotted portions of the molecules. This difference is better seen in the inset where the main histogram was normalized to bring each histogram bar to one. The normalized histogram clearly shows that if Topo IV were selective for the left-handed juxtapositions with the $\theta$ angle of about $-40^\circ$, then nearly all passages would result in unknotting and there would be practically no undesired passages causing partial relaxation of supercoiled DNA molecules.

Differences between the bending angle of potential G segments in knotted and interwound portions of supercoiled DNA molecules

Earlier observations of Topo IV bound to DNA (21) and earlier modeling studies of non-supercoiled knotted DNA molecules have suggested that Topo IV can unknot non-supercoiled DNA efficiently by acting preferentially at the juxtapositions involving strongly bent DNA regions (21,22,24,43). However, in bacteria DNA molecules are supercoiled. Therefore, it is important to check whether selection of bent DNA regions can also favor unknotting in supercoiled DNA molecules. Molecules shown in Figure 3 give an impression that the DNA portion in the knotted regions is more highly bent than in the plectonemic regions. A similar observation was made earlier (42) but it was not determined whether strand passages within DNA-DNA jux-
tapproaches one. We can see that the proportion of juxtapositions that result in unknotting upon intersegmental passage increases as the α₁ angle decreases. Therefore, by acting preferentially on juxtapositions with strongly bent DNA, Topo IV would be expected to increase, approximately 10-fold, its chances to unknot supercoiled DNA molecules when compared to acting on the juxtapositions with a weakly bent G segment. However, even for highly bent G segments, the probability of unknotting does not exceed 60%. In contrast, in juxtapositions with an angle of about −40°, the probability of unknotting approaches one (see inset in Figure 4A).

The combined effects of two different geometric characteristics of DNA-DNA juxtapositions on the topological outcome of the strand passage

The geometrical selection of potential sites of action by topoisomerases is likely to be simultaneously affected by several geometric characteristics of DNA–DNA juxtapositions. In general, the configurations of the juxtapositions that best fit the DNA binding sites of the enzyme will be recognized with the highest probability. By combining several geometric characteristics of DNA–DNA juxtapositions, the topoisomerase’s specificity for unknotting can be enhanced. However, it also is possible that if one selection criteria is fulfilled, the other may become less important. Figure 4C shows the complex effect of combining just two characteristics, i.e. the crossing angle θ and the bending angle α₁, on the topological outcome of strand passage. The color code in Figure 4C (see the scale) indicates the probability with which juxtapositions with a given combination of the θ and α₁ angles result in unknotting the supercoiled knotted DNA molecules. We can see that a relatively large region of the configuration space that predisposes DNA–DNA juxtapositions for very efficient unknotting lies in the favorable intersection of θ angles ranging from −20° to −60° and α₁ angles ranging from 100° to 180°. However, very strong bending of the G segment increases the probability of unknotting regardless of the θ angle.

The structure of negatively supercoiled postreplicative DNA catenanes

As early as the 1980s, it was observed that just after replication, circular DNA molecules form multiply interlinked right-handed catenanes (8,9). This interlinking is a consequence of the right-handed helical structure of DNA (8). Of course, freshly replicated DNA molecules need to be

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**Figure 4.** Geometrical and topological characterization of juxtapositions in the modeled supercoiled DNA molecules forming right-handed trefoil knots. (A) Main part: Raw data, a stacked histogram in which the heights of the histogram bars correspond to the total number of juxtapositions observed in the analyzed configurations falling within a given range of the θ angle. The colors indicate the fractions of the juxtapositions (within a given θ range) that, upon intersegmental passages, result either in unknotting (blue) or maintaining the original knot type (green). Juxtapositions that result in unknotting involve knotted portions of the molecules, whereas juxtapositions that maintain the original knot type are located within the interwound portions. Inset: The normalized histogram from the main part shows more directly that the fraction of juxtapositions that result in unknotting upon intersegmental passage increases as the α₁ angle decreases. Therefore, by acting preferentially on juxtapositions with strongly bent DNA, Topo IV would be expected to increase, approximately 10-fold, its chances to unknot supercoiled DNA molecules when compared to acting on the juxtapositions with a weakly bent G segment. However, even for highly bent G segments, the probability of unknotting does not exceed 60%. In contrast, in juxtapositions with an angle of about −40°, the probability of unknotting approaches one (see inset in Figure 4A).
decatenated rapidly and several different DNA topoisomerases participate in this process. From studies of bacterial systems, it is known that when freshly replicated DNA molecules are not yet covalently closed, and contain nicks and gaps, their decatenation can be performed by topoisomerase III (44). Topo III is a type I DNA topoisomerase that preferentially performs passages between single and double stranded regions of the DNA. The activity of DNA gyrase is needed for efficient decatenation by Topo III (45) and it was suggested that the action of DNA gyrase on nicked and gapped DNA can result in pushing the interlinks toward nicks and gaps (46). Once both freshly replicated molecules become covalently closed and supercoiled, Topo IV is responsible for their further decatenation (1). To investigate how Topo IV can efficiently decatenate supercoiled DNA molecules without relaxing their supercoiling, we performed simulations of multiply catenated supercoiled DNA molecules. The simulated molecules are interlinked with each other five times and have the supercoiling density typical for bacterial plasmids.

Figure 5 shows two examples of equilibrated configurations of the simulated catenanes. Similar to the confinement and localization of the knotted portions in supercoiled molecules (see Figure 3), the catenated portions are confined and tend to localize at the apical loops of supercoiled DNA molecules.

Negative crossing angles $\theta$ are strongly enriched in the catenated portions of negatively supercoiled, catenated DNA molecules

The configurations of negatively supercoiled, catenated DNA molecules shown in Figure 5 give the impression that the crossing angles in the catenated regions and in the plectonemic interwound portions of the molecules are quite similar to each other. To analyze in more detail possible differences between the $\theta$ angles of juxtapositions formed in the catenated or in interwound portions of the molecules, we characterized the geometry and the topological outcome of strand passages occurring at the given juxtapositions. As is the case in knotted DNA molecules, one can infer the location of a given juxtaposition from the topological outcome of passages between juxtaposed segments in that juxtaposition. If the passage in a given juxtaposition leads to a decrease in the catenation number of catenated molecules, the juxtaposition in question involves segments belonging to the region of catenation. On the other hand, strand passages in the juxtapositions located in the interwound portions result in molecules keeping the original catenation number. The raw data, stacked histogram in Figure 6A shows the total number of juxtapositions within a given interval of $\theta$ angles and also tells how many of them lie in the interwound and catenated portions of the molecules (indicated on the histogram with green and blue color, respectively). Upon normalization of each histogram bar to one (see the inset), we see that among the relatively rare juxtapositions with negative $\theta$ angle, there is strong domination of juxtapositions formed in the catenated portions since passages occurring within them lead to a decrease of the catenation number (topological change from the catenane $10^2$ to $8^2$). This analysis shows that if Topo IV were selectively acting on juxtapositions with the $\theta$ angles ranging between $-40^\circ$ and $-60^\circ$, the great majority of the passages would result in progressive decatenation of the DNA molecules. It is interesting to note that a similar range of $\theta$ angles also was optimal for the unknotted of supercoiled DNA molecules forming right-handed trefoil knots (see Figure 4).

Preferential interaction with bent DNA regions can aid Topo IV in progressive decatenation of catenated supercoiled DNA molecules

The raw data histogram shown in Figure 6B shows that the most frequently observed $\alpha_1$ bending angles range from $140^\circ$ to $160^\circ$ and that this is true for both the juxtapositions in the catenated and the interwound portions of the modeled DNA molecules. The normalized histogram (inset) reveals though that by selecting juxtapositions with more highly bent G segment (i.e. with a smaller $\alpha_1$ angle), Topo IV can increase the fraction of passages leading to progressive decatenation. However, the effect is only moderate and other additional selection criteria would be needed to ensure highly specific decatenation.

Figure 6C shows the combined effect of the crossing angle $\theta$ and the bending angle $\alpha_1$ on the topological outcome of passages occurring in the juxtapositions within different angle ranges. The color scale indicates the probability that passages involving juxtapositions with a given geometric char-
acteration results in a decrease of their catenation number from 5 to 4 and passage from catenanes with the topological notation $10^2$ to $8^2$. We see that the $\theta$ angles ranging from $-40^\circ$ to $-60^\circ$ strongly predispose juxtapositions for progressive decatenation.

The location of juxtapositions that can be selected by Topo IV for decatenation-specific passages

As is shown in Figure 6A, only a very small fraction of juxtapositions in catenated, supercoiled DNA molecules have left-handed crossings with the $\theta$ angle ranging from $-40^\circ$ to $-60^\circ$. These juxtapositions are highly enriched in the catenated portions of the molecules and thus Topo IV’s specificity for juxtapositions with such a geometry could result in targeted decatenation. In fact, Topo IV is likely to have such specificity since left-handed juxtapositions within this range of $\theta$ angle dominate in positively supercoiled DNA molecules that are preferentially recognized by Topo IV (4,15). To understand better how Topo IV can specifically decatenate negatively supercoiled right-handed catenanes, we investigated where the left-handed crossings with $\theta$ angles ranging from $-40^\circ$ to $-60^\circ$ are localized in these molecules. Our analysis reveals that these juxtapositions are preferentially located in the region where plectonemic portions of catenated molecules join the catenated portions (see Figure 7). As illustrated in Figure 7, one of the crossings formed in this region is left-handed, whereas all other crossings in the configuration are right-handed. In addition, this crossing is between segments belonging to two freshly replicated daughter molecules. Therefore, Topo IV, which shows a strong preference for left-handed crossings (4), can nevertheless very specifically decatenate right-handed catenanes formed by supercoiled DNA molecules since one specific crossing in these catenanes is frequently left-handed.

DISCUSSION

It has been a puzzle as to how Topo IV, which shows selectivity for left-handed crossings, can efficiently decatenate multiply interlinked right-handed catenanes. A partial answer to this question was provided by Stone et al. who measured the activity of Topo IV on right- and left-handed DNA braids (4). These measurements, combined with numerical simulations of DNA braids, led the authors to propose that Topo IV can act only on left-handed crossings, such as those omnipresent in positively supercoiled DNA. In addition, Stone et al. proposed that the left-handed crossings also can form in right handed catenanes provided that the catenanes have a low catenation number (4). A lack of supercoiling combined with a low catenation number give these catenated molecules enough structural freedom to be driven by thermal motion to occasionally form left-handed crossings. However, Stone et al. did not consider that freshly replicated molecules can have high catenation number and that these catenated DNA molecules get supercoiled before their decatenation (47). Therefore, it is important to investigate whether these additional structural constraints can prevent supercoiled DNA molecules with relatively high catenation number from forming left-handed crossings suitable for Topo IV-mediated decatenation.
crossings with \( \theta \) angles typical for positively supercoiled DNA molecules appear very rarely in negatively supercoiled DNA, which explains how negatively supercoiled DNA can be protected from the Topo IV action. Without such protection, the Topo IV action would result in the continuous relaxation of the negative supercoiling.

Our simulation results support the original proposal of Stone et al. that 'Topo IV has a strong preference for left-handed juxtapositions with the \( \theta \) angle ranging from \(-30^\circ\) to \(-60^\circ\) (4). However, the 2003 proposal of Stone et al. was challenged in 2009 by Neuman et al. (6). Using magnetic tweezers to wrap two DNA molecules around each other at different crossing angles, Neuman et al. concluded that the optimal crossing angle for Topo IV to act is very close to the achiral crossing, where two contacting segments are nearly perpendicular to each other (\( \theta \approx 0^\circ \)) (6). However, the experimental setup used by Neuman et al. introduced a coupling between the \( \theta \) angle and the \( \alpha_3 \) angle of enforced juxtapositions. Juxtapositions, where the approaching segments were nearly perpendicular to each other were also most likely to be strongly bent. As discussed in the introduction, Topo IV has high affinity to strongly bent sites. Therefore, we believe that the preference of Topo IV for juxtapositions with nearly perpendicular crossing angle \( \theta \) observed in experiments by Neuman et al. was likely the result of the high affinity of Topo IV to strongly bent DNA and did not correctly reflect the enzyme’s preference for a given \( \theta \) angle. Interestingly, our simulations show that juxtapositions involving strongly bent G segments predispose knotted molecules for unknotting and catenated molecules for decatenation even when the \( \theta \) angle is not in the \(-40^\circ\) to \(-60^\circ\) range.

The geometric selection of juxtapositions by Topo IV and other topoisomerases is likely to depend on several geometric characteristics of these juxtapositions. Earlier studies have shown that by selecting strongly bent DNA, Topo IV can efficiently unknot torsionally relaxed DNA molecules (21,22,24,43). Our simulation results show that the selection of \( \theta \) angles in the range \(-40^\circ\) to \(-60^\circ\) aided by the preferential interaction with bent DNA sites can explain how Topo IV efficiently unknots and also decatenates negatively supercoiled DNA molecules without causing their torsional relaxation. In crystals where Topo IV interacts with G segments, these DNA regions are strongly bent (25). Although there are not yet any solved crystals in which the G and T segments are bound by Topo IV, our results lead us to anticipate that the \( \theta \) angle between the G and T segments in such crystals will be between \(-40^\circ\) and \(-60^\circ\).

Our study focused on the situation where the replication is already finished and both freshly replicated daughter DNA molecules, forming catenanes, are covalently closed and supercoiled. However, when the replication is not yet completed, the freshly replicated portions of circular DNA molecules form the so-called precatenanes (48). Since actively replicated DNA molecules accumulate positive supercoiling the precatenanes are expected to wind around each other in a right-handed fashion (3,48). As long as the replication is not finished, the freshly replicated portions forming precatenanes contain discontinuities in freshly synthesized strands and this precludes the precatenanes from accumulating torsional stress needed for supercoiling. For this

Our numerical simulations of negatively supercoiled, catenated DNA molecules with relatively high catenation number show that the catenated region is highly confined in such molecules and that this confinement limits the possibility of forming short regions with left-handed windings within this region. However, in the regions where the plectonemic portions join the catenated portion, there is one juxtaposition (crossing) that is left-handed (see the magnified view), whereas all of the other crossings in the presented configuration are right-handed (see insets and Figure 3 for schematics of right- and left-handed crossings).

Figure 7. An equilibrated configuration of supercoiled, catenated DNA molecules. Notice that in the region where the plectonemic portions join the catenated portion, there is one juxtaposition (crossing) that is left-handed (see the magnified view), whereas all of the other crossings in the region of catenation, one observes the formation of individual left-handed juxtapositions involving segments of two catenated circular molecules (see Figure 7). Interestingly, the formation of these left-handed juxtapositions in right-handed catenanes is the natural consequence of the specific structure of multiply interlinked catenanes where each of the two catenated molecules is negatively supercoiled (see Figure 7). These left-handed juxtapositions form in regions where the plectonemic portions join the catenated portion and have crossing angles \( \theta \) typical for positively supercoiled DNA. Importantly, left-handed
reason, it is unlikely that right-handed precatenanes will form left-handed crossings of the type that we observe in the modeled supercoiled catenanes. Topo IV would be, therefore, unlikely to decatenate these precatenanes. However, we showed recently that gyrase action on nonsupercoiled catenanes is likely to promote topoisomerase III-mediated decatenation occurring at nicks and single stranded gaps (46). Such an activity is independent of the handedness of the interwinding of the contacting DNA segments and thus should also be efficient in promoting decatenation of right-handed precatenanes.

The modeling studies presented here did not account for the fact that DNA in bacterial cells interacts with histone-like proteins such as H-NS and HU (49). As a consequence of this interaction, part of the supercoiling of bacterial DNA is constrained and the magnitude of supercoiling density in vivo is roughly twice lower than in the deproteinized DNA isolated from bacterial cells (50–53). We did not account for this decrease of the magnitude of supercoiling density in vivo. However, our earlier modeling studies showed that conditions of high crowding, characteristic for bacterial nucleoids, naturally compensate for this decreased magnitude of supercoiling in vivo (32).

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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