Requirements for DNA-Bridging Proteins to Act as Topological Barriers of the Bacterial Genome

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ABSTRACT Bacterial genomes have been shown to be partitioned into several-kilobase-long chromosomal domains that are topologically independent from each other, meaning that change of DNA superhelicity in one domain does not propagate to neighbors. Both in vivo and in vitro experiments have been performed to question the nature of the topological barriers at play, leading to several predictions on possible molecular actors. Here, we address the question of topological barriers using polymer models of supercoiled DNA chains that are constrained such as to mimic the action of predicted molecular actors. More specifically, we determine under which conditions DNA-bridging proteins may act as topological barriers. To this end, we developed a coarse-grained bead-and-spring model and investigated its properties through Brownian dynamics simulations. As a result, we find that DNA-bridging proteins must exert rather strong constraints on their binding sites; they must block the diffusion of the excess of twist through the two binding sites on the DNA molecule and, simultaneously, prevent the rotation of one DNA segment relative to the other one. Importantly, not all DNA-bridging proteins satisfy this second condition. For example, single bridges formed by proteins that bind DNA nonspecifically, like H-NS dimers, are expected to fail with this respect. Our findings might also explain, in the case of specific DNA-bridging proteins like LacI, why multiple bridges are required to create stable independent topological domains. Strikingly, when the relative rotation of the DNA segments is not prevented, relaxation results in complex intrication of the two domains. Moreover, although the value of the torsional stress in each domain may vary, their differential is preserved. Our work also predicts that nucleoid-associated proteins known to wrap DNA must form higher protein-DNA complexes to efficiently work as topological barriers.

INTRODUCTION

The genetic information of most bacteria is encoded in a circular DNA molecule comprising up to several millions of basepairs (bp). Such closed molecules are topologically constrained because they lack the free ends capable of rotating and releasing the torsional stress (1). In vivo alterations of the torsional state of circular DNA is mediated by the recruitment of enzymes called topoisomerases, which can either increase or decrease the twist of the double helix by opening transiently one or two strands (1). The net result of the action of all types of topoisomerases is that the genomic DNA of most bacteria is significantly undertwisted (negatively supercoiled) and winds about itself to transfer part of the torsional stress to the bending degrees of freedom, thereby forming plectonemes (1). If the DNA were not subject to any additional constraint beyond closure, then one single nick of one strand or one single break of the two strands would suffice to release the torsional stress of the full molecule. It has, however, long been known that at least several tens of nicks are required to achieve this goal in Escherichia coli (2,3). This indicates that there exist

SIGNIFICANCE Although the question of independent topological domains in bacterial genomes has been addressed for nearly two decades, the nature of topological barriers is still elusive. The most plausible models for the formation of topological barriers are actively transcribing RNAP and the formation of DNA loops by certain nucleoid proteins. In this work, we focus on this latter mechanism and establish under which conditions DNA-bridging proteins may serve as topological barriers. We show that, contrary to popular belief, formation of a loop is not sufficient and that proteins must additionally block the diffusion of twist and prevent the relative rotation of DNA segments. These findings set strong theoretical limits on the ability of bridging proteins to create topologically independent genomic domains.
barriers that block the diffusion of the torsional stress along the DNA molecule and organize the chromosome of *E. coli* into many independent topological domains whose torsional state is not affected by the relaxation of other domains. It is currently estimated that the genomic DNA of bacteria like *E. coli* is composed of several hundreds of different topological domains with variable size (average size ~10 kbp) and position (4). More generally, the partitioning of the bacterial chromosome into ~10 kbp independent topological domains is believed to hold in most bacteria (5) and to be related, in part, to the insulation of fundamental coexpression units that are neighbors along the genome (6).

The work reported here deals with the topological barriers that make this partitioning possible, that is, the barriers that block the diffusion of torsional stress along the DNA molecule and are responsible for its division into independent topological domains. Although this question has now been addressed for nearly two decades (7–9), the mechanisms underlying the formation of topological barriers remain mostly elusive. The most plausible models for the formation of topological barriers are 1) actively transcribing RNA polymerases (RNAPs), which generate both positive and negative supercoils (10) and may consequently block the displacement and dissipation of plectonemes (11,12); and 2) the formation of DNA loops by certain nucleoid proteins that may serve as topological barriers (13–16). In this work, we focus on this latter mechanism and establish under which conditions DNA-bridging proteins may serve as topological barriers.

Several DNA-bridging proteins have been studied in some detail, including transcription regulators like the LacI repressor (17), H-NS (18) and H-NS-like proteins (19), Lsr2 (20), Fis (21), and Lrp (22). In their functional form, all of these proteins have at least two independent DNA-binding domains so that they can interact with two DNA duplexes simultaneously and form a bridge between two sites that are widely separated from the genomic point of view. Most of these proteins bind nonspecifically but with high affinity to the DNA molecule. However, among all these DNA-bridging proteins, LacI is, to date, the only one that has been proved to work as a topological barrier (15,23,24).

The guiding line of this study is, consequently, the experimental demonstration in (15) that the binding of a DNA-binding protein to its recognition sites in two different locations on a supercoiled DNA molecule can confine free supercoils to a defined region and divide the DNA molecule into two distinct topological domains (15). To gain more detailed information on this mechanism, we determined the minimal physical properties that must be added to a standard model of circular DNA to reproduce the results described in (15). The starting model for this study was similar to those we proposed recently to investigate facilitated diffusion (25–27), the interactions of DNA and H-NS nucleoid proteins (28–30), the formation of the bacterial nucleoid (31–35), and the interplay of DNA demixing and supercoiling in compacting the nucleoid (36).

The results presented in this article reveal that the formation of DNA loops is by no means sufficient to create topologically independent domains and that DNA-bridging proteins must exert rather strong constraints on their binding sites to act as topological barriers; they must block the diffusion of the excess of twist through both binding sites and must additionally block the rotation of one DNA segment relative to the other one.

**METHODS**

The coarse-grained bead-and-spring model developed for this study is described in detail in Supporting Materials and Methods, Model and Simulations. In brief, bacterial DNA is modeled, as in (30,36), as circular chains of *n* beads with a radius *a* = 10 nm separated at equilibrium by a distance *l* = 2.5 nm, where two beads represent 15 DNA bp. Simulations were performed with unconstrained circular chains of length *n* = 600, equivalent to 4500 bp, which represent the plasmids used in (15). DNA beads interact through stretching, bending, torsional, and electrostatic terms. The bending rigidity constant (*g = 20k_bT*) was chosen so that the model reproduces the known persistence length of double-stranded DNA (*ξ = 50 nm*).

As in (31,36), the torsional energy term was borrowed from (37) and requires the introduction of a body-fixed frame (*u_α, f_α, v_α*), where *u_α* denotes the unit vector pointing from bead *k* to bead *k + 1*. The torsional rigidity opposing rotation of (*u_α, f_α, v_α*) around *u_α* (*τ = 25 k_bT*) was adjusted so that at equilibrium the writhe contribution accounts for ~70% of the linking number difference (38), as is illustrated in Fig. S1. The values of the bending and torsional rigidities are close together, in agreement with experimental results (38). It may be worth emphasizing that introduction of the body-fixed frame (*u_α, f_α, v_α*) is crucial for correctly modeling the torsion of double-stranded DNA with a “single-stranded” bead-and-spring model. This procedure is quite realistic and has already proved very useful for studying, for example, the interplay of DNA demixing and supercoiling in nucleoid compaction (36), the buckling transition in double-stranded DNA and RNA (39), the influence of nucleoid-associated proteins on DNA supercoiling (40), DNA supercoil relaxation (41), the competition between B-Z and B-L transitions in a single DNA molecule (42), the relaxation of DNA supercoils by topoisomerase IB (43), the sequence-dependent duplex denaturation in superhelical DNA (44), transcription-driven twin supercoiling of DNA loops (45), site juxtaposition in supercoiled DNA (46), and salt effects on the structure and internal dynamics of superhelical DNA (47).

Electrostatic repulsion between DNA beads is written as a sum of Debye–Hückel terms, which depend on effective electrostatic charges placed at the center of each bead. The values of these charges (*q = −3.52e*, where *e* is the absolute charge of the electron) were derived from the net linear charge density along a DNA molecule immersed in a buffer with monovalent cations according to Manning’s counterion condensation theory (48,49). The value of the Debye length (*ξ_D = 1.07 nm*) corresponds to a concentration of monovalent salt of 100 mM, which is the value that is generally assumed for the cytoplasm of bacterial cells.

The torsional state of the DNA chain is characterized by the value of the superhelical density *σ = ΔLk/Lk_0*, where the linking number *Lk_0 = 7.5n_l/10.5 ~ 429* is the ratio of the number of base pairs of the DNA chain and the mean number of base pairs per turn of the torsionally relaxed double helix, and the linking number difference *ΔLk = ΔTw + Wr* is the sum of the excess of twist (*ΔTw*) and the writhe (*Wr*), which quantifies the winding of the DNA axis around itself. *ΔLk* and *σ* are constants of the motion, whereas *ΔTw* and *Wr* fluctuate under the influence of thermal noise and external constraints. Bacterial DNA being negatively supercoiled (underwound), *ΔLk* and *σ* are negative in this work. Investigated values of *σ* range from
–0.033 to –0.121, which is approximately from the effective value for protein-bound DNA in living E. coli cells (σ ≈ –0.025 (50)) to about twice the average value for naked DNA in vitro (σ ≈ –0.06 (51)). σ ≈ –0.12 is the maximal value generated by DNA gyrase and RNAP (52–54) and corresponds to a range of superhelicity values in which hyperplectonemes, a higher-order DNA organization, have been observed (55,56). The properties of the model were determined by integrating numerically overdamped Langevin equations with time steps Δt = 10 ps. Temperature T was assumed to be 298 K throughout the study.

RESULTS AND DISCUSSION

As a starting point, we recall the experimental results in (15), in which the torsional relaxation properties of 4100 bp plasmids with initial superhelical density σ = –0.06 (ΔLk ≈ –24) were thoroughly investigated. Of special interest to us were the experiments performed with plasmids containing tandem copies of the sequence-specific DNA-binding protein LacI at two different locations such that the plasmid was divided into two stable loops of respective lengths of 2900 and 1200 bp upon binding of LacI. The 1200 bp region moreover contained the recognition site for one nicking enzyme (Nt.BbvC1) and the 2900 bp region the recognition site for a second nicking enzyme (Nb.BtsI). In the absence of LacI, the addition of Nt.BbvC1 alone or Nb.BtsI alone was sufficient to release the full torsional stress of the plasmid. In contrast, in the presence of LacI, the addition of Nt.BbvC1 alone removed only the seven negative supercoils of the 2900 bp region (15). Simultaneous addition of both enzymes was required to relax the full plasmid (15). This experiment demonstrates very clearly that pairs of LacI bridges block the diffusion of torsional stress and divide the plasmids into two independent topological domains.

We report below on our efforts to determine the minimal set of mandatory properties that allow molecular bridges to act as topological barriers and block the diffusion of torsional stress in out-of-equilibrium supercoiled chains.

Preparation of torsionally out-of-equilibrium circular chains

Topological barriers are objects that block the diffusion of torsional stress and are able to maintain different torsional conditions on their left and right sides. Characterization of the underlying mechanism requires prior understanding of how torsional stress diffuses in out-of-equilibrium supercoiled chains. To prepare such chains, we first let circular chains with n = 600 beads and different values of ΔLk equilibrate for ~100 ms. ΔLk was set to ΔLk0 = −14, −20, −26, −34, −40, −46, and −52, corresponding to σ0 ≈ −0.033, −0.047, −0.061, −0.079, −0.093, −0.107, and −0.121, respectively. Equilibrated chains were then formally divided into two moieties of equal length on both sides of beads α (1 ≤ α ≤ n/2) and β = α + n/2, which were chosen so that the distance between their centers is smaller than 10 nm. This step is illustrated in Fig. 1 a, which shows an equilibrated chain with ΔLk0 = −40 (σ0 ≈ −0.093), where beads α ≤ k ≤ β are colored in green and the other ones in red. The torsional stress in segment α ≤ k ≤ β was then artificially released by rotating the internal basis (ui, fi, vj) around ui by an angle 2π(k − α) ΔLk0/n for α ≤ k ≤ β, thereby increasing the twist in this segment by an amount equal to ΔLk0/2. Finally, the segment α ≤ k ≤ β was equilibrated again for 5 ms, whereas the other moiety was kept frozen. As illustrated in Fig. 1 b, this led to torsionally out-of-equilibrium circular chains with one torsionally relaxed moiety such that ΔLk1 ≡ ΔLk1 = W1 = 0 and one torsionally stressed moiety such that ΔLk2 ≡ ΔLk2 = 0.3 ΔLk2, and W2 = 0.7 ΔLk2. The ΔLk1 and W1, j = 1, 2 are the partial excess of twist and writhe for each moiety, and the ΔLk = ΔLk1 + W1 are the partial linking number differences. These quantities are introduced in more detail in Supporting Materials and Methods, Model and Simulations, and satisfy

FIGURE 1 (a) Representative conformation of an equilibrated circular chain with linking number difference ΔLk = −40 (σ0 ≈ −0.093). The chain is separated formally into two moieties of equal length, which are colored in green and red, respectively. The black arrows locate beads α and β, which separate the two moieties. (b) Representative snapshot of the same chain after increasing the twist in the green segment by ΔLk/2 and letting this segment equilibrate again for 5 ms while keeping the red segment frozen. This is an out-of-equilibrium conformation. (c) Representative snapshot obtained upon relaxation of the out-of-equilibrium conformation shown in (b). The linking number difference of the equilibrated chain is ΔLk/2 = −20 (σ0 ≈ −0.047). To see this figure in color, go online.
\[ \Delta T_w = \Delta T_{w1} + \Delta T_{w2} \]
\[ W_t = W_{t1} + W_{t2} + W_{t12} \]
\[ \Delta L_k = \Delta L_{k1} + \Delta L_{k2} + W_{t12}, \]

where \( W_{t12} \) quantifies the winding of one moiety of the chain around the other one that is, loosely speaking, their intrication. The out-of-equilibrium chains prepared as described above satisfy \( W_{t12} \approx 0 \).

The main purpose of this work is to understand how DNA-bridging proteins like LacI (15,23,24) can maintain such an unbalance when the whole chain is relaxed. Because DNA-bridging proteins form DNA loops by dynamically cross-linking widely separated DNA sites, the question amounts here to determine the constraints that must be imposed to beads \( \alpha \) and \( \beta \) to divide the circular DNA chain into two topologically independent loops. Note that DNA-bridging proteins are not introduced explicitly in the model. Rather, constraints on beads \( \alpha \) and \( \beta \) are meant to model the mechanisms that allow the proteins to act as topological barriers.

**Twist equilibrates much more rapidly than writhe**

To understand in more detail how the diffusion of torsional stress proceeds, the out-of-equilibrium circular chains prepared as described above were allowed to relax without constraint, and all relevant quantities were monitored. From a practical point of view, four different initial out-of-equilibrium conformations were prepared for 1 ms for each initial conformation. The evolution of \( \Delta L_{k1}, \Delta T_{w1}, \) and \( W_t (j = 1, 2) \) was then averaged over these 100 trajectories. In the absence of any constraint on beads \( \alpha \) and \( \beta \), all conformations obtained at the end of the 1 ms integration time satisfy \( \Delta L_{k1} \approx \Delta L_{k2} \) and \( \Delta T_{w1} \approx \Delta T_{w2} \), meaning that torsional stress is distributed homogeneously in the whole chain. Information on the dynamics of the relaxation is provided by the examination of the time evolution of \( \Delta T_{w2} - \Delta T_{w1} \) and \( W_{t2} - W_{t1} \). As illustrated in Fig. S2 for an initial conformation with \( \Delta L_{k1} = -26 (\sigma^0 \approx -0.061) \), both quantities decay almost exponentially:

\[ \Delta T_{w2} - \Delta T_{w1} \approx b \exp \left( \frac{-t}{\tau_{tw}} \right) \]
\[ W_{t2} - W_{t1} \approx A + B \exp \left( \frac{-t}{\tau_{tw}} \right), \]

where \( \tau_{tw} \) and \( \tau_{w} \) are the characteristic time constants for the equilibration of the excess of twist and the writhe, respectively. Let us stress here that we are not aware of any reason why \( \Delta T_{w2} - \Delta T_{w1} \) and \( W_{t2} - W_{t1} \) should decay according to an exponential law, that is according to a process involving a single timescale, although such exponential decays might be rationalized by analyzing the relaxation dynamics of the DNA chain as a twist diffusion process (57). Equation 2 should thus be essentially considered here as a convenient means to extract and compare characteristic equilibration time constants for the different models investigated in this work, that is the circular DNA chain without constraints on beads \( \alpha \) and \( \beta \) discussed in this subsection and the three different models of DNA-bridging proteins discussed in the following subsections. The evolution of \( \tau_{tw} \) and \( \tau_{w} \) as a function of \(-\sigma^0\) is shown in Fig. 2. Each point in this figure was obtained from a fit of Eq. 2 against the time evolution of \( \Delta T_{w2} - \Delta T_{w1} \) and \( W_{t2} - W_{t1} \) for a different initial conformation. This figure indicates that \( \tau_{tw} \) increases slightly with \(-\sigma^0\) but remains comprised between 1 and 2 \( \mu s \) for all investigated values of \(-\sigma^0\). Equilibration of the excess of twist between the two moieties of the circular chain is consequently rather fast, the reason being that it involves only some rotation of the chain around its own axis—more precisely, rotation of the internal basis.
(\textbf{u}_k, \mathbf{f}_k, \mathbf{v}_k) around \textbf{u}_k for all \( k \). In contrast, Fig. 2 indicates that \( \tau_{\text{wr}} \) is typically one to two orders of magnitude larger than \( \tau_{\text{rw}} \), which means that the equilibration of the writhe is much slower than the equilibration of the excess of twist. The reason for this longer time constant is that the equilibration of the writhe requires global rearrangements of the whole chain, as can be checked in Fig. 1, instead of its mere rotation around its own axis. Moreover, the larger the difference in torsional stress between the two moieties, the larger the torsional force and torque exerted by each moiety on the other one, and the faster the motion of the beads involved in the rearrangement. This is probably the reason why \( \tau_{\text{wr}} \) decreases significantly with increasing \( |\sigma^0| \), from \( \sim250 \ \mu s \) at \( \sigma^0 \approx -0.033 \) (in the same range as reported in (41)) down to \( \sim40 \ \mu s \) at \( \sigma^0 \approx -0.121 \). Finally, the values of \( \tau_{\text{wr}} \) vary more broadly from one initial conformation to the other one for smaller values of \( |\sigma^0| \) than for larger ones, as can be checked in Fig. 2.

At equilibrium, the linking number difference decomposes into \( \sim70\% \) of writhe and \( \sim30\% \) of excess of twist (see Fig. S1), but equilibration of the excess of twist is much faster than the equilibration of the writhe. After a couple of microseconds, the linking number difference in the initially relaxed moiety of the chain is therefore composed of \( 0.5 \times 0.3 \times 0.5 \times \Delta L k^0 = 0.075 \Delta L k^0 \) excess of twist and no writhe (0% writhe and 100% excess of twist), whereas in the initially stressed moiety it decomposes into \( 0.7 \times 0.5 \times \Delta L k^0 = 0.35 \Delta L k^0 \) of writhe and \( 0.5 \times 0.3 \times 0.5 \times \Delta L k^0 = 0.075 \Delta L k^0 \) excess of twist (82% writhe and 18% excess of twist). As a consequence, during the most part of the equilibration process, the initially torsionally relaxed moiety of the chain has not enough writhe compared with the excess of twist, whereas the initially torsionally stressed moiety has too much writhe compared with the excess of twist. The need to equilibrate the writhe/twist ratio inside each moiety of the DNA chain may in turn accelerate the equilibration of writhe in the whole chain by favoring diffusion of the writhe from the initially stressed moiety to the initially relaxed one.

Several comments are in order here. First, the faster equilibration of twist compared with writhe for closed molecules or filaments with both torsional and bending energy (and comparable torsional and bending force constants) is well known in the field of DNA biophysics. Yet, there are relatively few studies that discuss this point explicitly and provide estimates for the characteristic relaxation timescales. We note, however, that the theoretical study in (58) already pointed out that the twist and writhe equilibrate on very different timescales. Second, it is emphasized that timescales deduced from coarse-grained models always underestimate real timescales by two to three orders of magnitudes because the coarse-grained models neglect many details that systematically slow down the dynamics. This point should, of course, be kept in mind when analyzing the results presented here and comparing them with experimental results. Third, we tested the importance of rotational noise (the last term in the right-hand side of Eq. S9) by launching simulations in which this term was not taken into account. Characteristic equilibration times obtained as described above are shown in Fig. S3. A comparison of Figs. 2 and S3 indicates that results obtained with and without rotational noise are identical within computational uncertainties. Rotational noise was consequently ignored in all subsequent simulations.

### Closing a DNA loop does not suffice to create an independent topological domain

Based on recent experimental results (12–16), one may wonder whether DNA loops are ipso facto independent topological domains or whether some additional, more subtle properties of DNA-bridging proteins are required to achieve this goal. We emphasize that the word “loop” is used here according to its biological meaning and not the geometrical one. More precisely, it does not refer to a closed curve but rather to a conformation of the DNA chain in which two DNA sites that are widely separated from the genomic point of view are maintained close to each other by some other macromolecule, so that the segment of DNA located between these two sites looks like a loop when it is observed from a long distance or projected on a plane. However, the chain is not closed, and torsional stress cannot diffuse directly from one extremity of the loop to the other one through the bridging macromolecule.

To ascertain this point, we launched a series of simulations with the crudest model for DNA-bridging proteins, namely a simple bond between beads \( \alpha \) and \( \beta \). The potential energy of the system was written as \( E_{\text{pot}} + E_{\text{BP}} \), where \( E_{\text{pot}} \) is the energy of the naked circular DNA chain described in Eqs. S1–S5 and \( E_{\text{BP}} \) is an additional term that models the action of DNA-bridging proteins. For the simplest model, we used

\[
E_{\text{BP}} = \frac{h}{2}(d_{\text{ab}} - d_{\text{ab}}^0)^2, \tag{3}
\]

where \( d_{\text{ab}} \) is the distance between the centers of beads \( \alpha \) and \( \beta \), and \( d_{\text{ab}}^0 = 4 \) nm. For the sake of simplicity, we used the same value of the stretching rigidity as for the DNA chain, that is \( h = 100k_B Tl_0^2 \) (see Supporting Materials and Methods, Model and Simulation). The value \( d_{\text{ab}}^0 = 4 \) nm is small enough to ensure that no other DNA segment crosses the segment between \( \alpha \) and \( \beta \), which is formally occupied by the DNA-bridging protein. The energetic barrier for any segment to cross this line is indeed about 32.8k_BT, which is virtually insuperable. For peace of mind, we nevertheless ran several simulations with \( d_{\text{ab}}^0 = 3 \) nm, corresponding to a barrier of about 60k_BT, for the second model discussed below and obtained exactly the same results as with \( d_{\text{ab}}^0 = 4 \) nm. This confirms that Eq. 3 describes an impenetrable \( \alpha \)-\( \beta \)-bond. The same initial conformations were used as in the preceding subsection, and
100 different relaxation trajectories were again integrated for 1 ms for each initial conformation. Characteristic equilibration times obtained as described above are shown in Fig. S4. Comparison of Fig. S4 with Figs. 2 and 3 indicates that the addition of a bond between beads \( \alpha \) and \( \beta \) affects significantly neither \( \tau_{r_\alpha} \) nor \( \tau_{r_\beta} \). Stated in other words, not all DNA-bridging proteins are topological barriers, or equivalently, not all DNA loops are independent topological domains. The ability of certain DNA-bridging proteins, like LacI, to act as topological barriers is consequently related to some property, which is subtler than the mere connection between two genetically widely separated DNA sites.

Still, it is worth noting that the \( \alpha\beta \)-bond has a significant effect on the time evolution of \( W_{12} \), as can be checked in Fig. S5. The reason is the following. To compute the partial writhes \( W_j \) (see Supporting Materials and Methods), one has to introduce two “virtual” closed chains, \( C_1 \) and \( C_2 \). \( C_1 \) is composed of beads \( \alpha \) to \( \beta - 1 \), plus the “unphysical” segment between beads \( \beta \) and \( \alpha \), whereas \( C_2 \) is composed of beads \( \beta \) to \( \alpha - 1 \), plus the same “unphysical” segment between beads \( \alpha \) and \( \beta \). When beads \( \alpha \) and \( \beta \) are bonded, they remain close to each other for \( t > 0 \), as indicated by the positions of the two arrows in Fig. 3 a, so that the unphysical segment between beads \( \alpha \) and \( \beta \) remains small, and \( C_1 \) is a good representation of the initially torsionally relaxed DNA loop (and \( W_1 \) is a measure of its writhes). \( C_2 \) is a good representation of the initially torsionally stressed DNA loop (and \( W_2 \) is a measure of its writhes), and \( W_{12} \) is a measure of the intrication of the two loops. In this case, \( W_{12} \) remains close to 0 at all times, as can be checked in Fig. S5 b. This indicates that, for bridged DNA, equilibration of the torsional stress is not accompanied by any significant intrication of the two circular moieties. In contrast, for the system without the \( \mathcal{E}_{BP} \) term, beads \( \alpha \) and \( \beta \) do not remain close to each other for \( t > 0 \), as indicated by the positions of the two arrows in Fig. 1 c, so that the unphysical segment between beads \( \alpha \) and \( \beta \) becomes arbitrarily large, and \( C_1 \), \( C_2 \), \( W_1 \), and \( W_{12} \) lose their physical meaning. In this case, \( W_{12} \) varies quite widely with changing DNA conformations, as is illustrated in Fig. S5 a for an initial conformation with \( \Delta L k^0 = -52 \) (\( \sigma^0 \approx -0.121 \)).

**Blocking diffusion of the twist is not sufficient to create independent topological domains**

The reason why forming a loop is not sufficient to create independent topological domains is of course that the bond between beads \( \alpha \) and \( \beta \) described in the previous subsection is not able to block the rapid diffusion of the excess of twist through beads \( \alpha \) and \( \beta \) and the subsequent equilibration of the writhes. To form independent topological domains, the diffusion of the excess of twist must absolutely be blocked. Stated in other words, rotation of the internal basis (\( \mathbf{u}_k, \mathbf{f}_k, \mathbf{v}_k \)) around \( \mathbf{u}_k \) must be forbidden at \( k = \alpha, \beta \). From the biological point of view, diffusion of the excess of twist is probably blocked because the involved proteins bind to their recognition site by inserting loops in the major or minor grooves of the DNA duplex. For example, it has been shown that H-NS dimers insert one C-terminal loop inside the minor grooves of double-stranded DNA (59), thereby hindering rotation of the DNA around its own axis at this location. From the modeling point of view, diffusion of the twist can be blocked by imposing that \( \mathbf{f}_k \) and \( \mathbf{f}_b \) remain perpendicular to \( \mathbf{r}_\alpha, \mathbf{r}_\beta - \mathbf{r}_\alpha \) at all times (\( \mathbf{f}_a, \mathbf{r}_\alpha, \mathbf{r}_\beta = \mathbf{f}_b, \mathbf{r}_\alpha, \mathbf{r}_\beta = 0 \)), which is achieved by computing the angles

\[
\dot{\delta}_k = -\tan^{-1}\left(\frac{\mathbf{f}_k, \mathbf{r}_\alpha, \mathbf{r}_\beta}{\mathbf{v}_k, \mathbf{r}_\alpha, \mathbf{r}_\beta}\right)
\]  

\( (k = \alpha, \beta) \), after each integration step and rotating \( \mathbf{f}_k \) and \( \mathbf{v}_k \) around \( \mathbf{u}_k \) by this angle \( \delta_k \).

\[
\mathbf{f}_k \rightarrow \cos \dot{\delta}_k \mathbf{f}_k + \sin \dot{\delta}_k \mathbf{v}_k
\]

\[
\mathbf{v}_k \rightarrow -\sin \dot{\delta}_k \mathbf{f}_k + \cos \dot{\delta}_k \mathbf{v}_k,
\]

for \( k = \alpha, \beta \). We note that the \( \dot{\delta}_k \) remain small, and the corrections in Eq. 5 consequently also remain small, only if \( \mathbf{r}_\alpha, \mathbf{r}_\beta \) is not perpendicular simultaneously to \( \mathbf{f}_k \) and \( \mathbf{v}_k \), that is if
vectors \( \mathbf{r}_{\alpha, \beta} \) and \( \mathbf{u}_k \ (k = \alpha, \beta) \) are not collinear. To prevent such collinearity, two bending terms were added to \( E_{\text{BP}} \), namely

\[
E_{\text{BP}} = \frac{\hbar}{2} \left( d_{ab} - d_{\alpha\beta}^0 \right)^2 + \frac{5g}{2} \left( \xi_{\alpha} - \frac{\pi}{2} \right)^2 + \frac{5g}{2} \left( \xi_{\beta} - \frac{\pi}{2} \right)^2,
\]

where \( \xi_{\alpha} \) denotes the angle formed by beads \( \beta, \alpha, \) and \( \alpha + 1 \) and \( \xi_{\beta} \) denotes the angle formed by beads \( \alpha, \beta, \) and \( \beta + 1 \). Note that the bending rigidity for these two angles was assumed to be five time larger than the bending rigidity of the DNA chain in order for \( \xi_{\alpha} \) and \( \xi_{\beta} \) to deviate only moderately from \( \pi/2 \). In this second model, the action of DNA-bridging proteins is consequently modeled by the potential energy terms in Eq. 6 plus the small a posteriori corrections in Eq. 5.

The relaxation dynamics for this second model is much slower than for the first one so that each trajectory had to be integrated for as long as 30 ms. Four different relaxation trajectories were integrated for each initial conformation, and four different initial conformations were again used for each value of \( \Delta L k^0 \). The typical time evolution of the twist and writhe for an initial conformation with \( \Delta L k = -20 (\sigma^0 = -0.047) \) is shown in Fig. 4. At the level of single trajectories, the \( W_r \), \( W_{r12} \), and \( \Delta L k_j \) actually relax through a succession of plateaus followed by abrupt jumps, as is illustrated in Fig. S6 for \( W_{r12} \), whereas the time evolution is, of course, smoother when these quantities are averaged over several trajectories, as in Fig. 4 b. It is stressed that this behavior has no precise physical meaning but arises instead from the mathematical definition of the total writhe \( W_r \) in Eq. S11 and its somewhat arbitrary partitioning into the sum of the two partial quantities \( W_r \) and \( W_{r12} \) in Eq. S17. In particular, \( W_r \) does not display any sharp jump along single trajectories.

As can be checked in Fig. 4, the second model of DNA-bridging proteins succeeds in preventing equilibration of the twist and writhe in the two moieties of the chain. Indeed, \( \Delta W_r = \Delta W_{r1} \) and \( W_r = W_{r1} \) remain nearly constant (and \( \Delta L k_2 = \Delta L k_1 \) remains rigorously constant) throughout the integration interval. Nonetheless, this model still fails to describe topological barriers, because \( W_{r1} \) and \( W_{r2} \) do evolve in parallel until the limit \( W_{r1} + W_{r2} = 0 \) is reached (see Fig. 4 b). According to Eq. 1, this limit is equivalent to \( W_{r12} = W_r \). The limit \( W_{r12} = W_r \) indicates very strong intrication of the two moieties. Indeed, examination of the final conformations, like the one with \( \Delta L k^0 = -40 (\sigma^0 = -0.093) \) shown in Fig. 3 b, reveals very compact structures, with the initially torsionally relaxed (green) moiety winding around the initially torsionally stressed (red) moiety. Clearly, the intrication of the two moieties is achieved by transforming part of the plectonemic supercoils of the red moiety into toroidal supercoils. Another general feature of the final conformations is that the portion of the red moiety not involved in the winding with the green moiety has a compact geometry that probably results from the superposition of plectonemic and toroidal supercoils.

More insight into the dynamics of the relaxation is gained by plotting the time evolution of \( 2 \langle W_{r12} \rangle / \Delta L k^0 \), where \( \Delta L k^0/2 \) is the final linking number difference of the circular chain. The result is shown in Fig. 5, where \( W_{r12} \) has been averaged over the 16 relaxation trajectories that have been computed for each value of \( \Delta L k^0 \). The four curves with \( |\Delta L k^0| \leq 34 (|\sigma^0| \leq 0.079) \) superimpose at short times, which indicates that the relaxation speed is proportional to the total amount of torsional stress. However, for longer times and/or larger values of the linking number difference, a slower regime sets in that is probably due to the resistance opposed by plectonemes against the reorganization of the chain and the intrication of the two moieties. Finally, we note that after full relaxation, the writhe contribution accounts for \( \sim 90\% \) of the linking number difference, which is significantly larger than the 70% contribution in the absence of domains (see Fig. S1).

In conclusion, this second model of DNA-bridging proteins indicates that blocking the diffusion of the excess of...
twist at the two DNA loci bound by the proteins is sufficient to maintain a constant difference between the linking number difference, excess of twist, and writhe of the two moieties but that it is not sufficient to insure true topological independence. Indeed, the DNA chain evolves toward very compact and intricated final conformations, in which the topology of each moiety is different from the initial one. The relaxation limit $W_{12} = W_t$ (or equivalently, $W_{1} + W_{2} = 0$) appears nontrivial and intriguing to us. We suspect that the explanation involves subtle considerations that should deserve further work.

**Topological barriers must block diffusion of the twist and the relative orientation of the DNA segments**

The question that remains to be answered is, consequently: what additional constraint must the DNA-bridging protein exert on its binding sites to preserve not only the difference in torsional stress between the two moieties of the DNA chain but rather the precise initial torsional stress in each moiety? The feeling that emerges upon examination of conformations like the one shown in Fig. 3 b is that the winding of the initially relaxed moiety around the initially stressed one probably requires the relative rotation of the two DNA segments around the $\alpha\beta$-bond and that no winding would be possible if this rotation were blocked. To check this conjecture, a third bending term was added to $E_{BP}$, namely

$$E_{BP} = \frac{h}{2} (d_{\alpha\beta} - d_{\alpha\beta0})^2 + \frac{5g}{2} (\xi_\alpha - \pi/2)^2 + \frac{5g}{2} (\xi_\beta - \pi/2)^2$$

$$+ \frac{5g}{2} (\psi_{\alpha,\beta} - \psi_{\alpha,\beta0})^2,$$  \hspace{1cm} (7)

where $\psi_{\alpha,\beta}$ denotes the angle between vectors $r_{\alpha} + 1 - r_{\alpha}$ and $r_{\beta} + 1 - r_{\beta}$ and $\psi_{\alpha,\beta0}$ is the value of $\psi_{\alpha,\beta}$ at time $t = 0$. The new term in the right-hand side of Eq. 7 prevents large fluctuations of the relative orientations of the two DNA segments that are bridged by the protein. In particular, it blocks the rotation around the $\alpha\beta$-bond of one segment with respect to the other one. In this third model, the action of DNA-bridging proteins is consequently modeled by the potential energy terms in Eq. 7 plus the small a posteriori corrections in Eq. 5.

The relaxation dynamics for this third model was investigated along the same lines as the second model. The typical time evolution of the twist and writhe for an initial conformation with $\Delta L_{K0} = -52$ ($\sigma^0 = -0.121$) is shown in Fig. S7. As can be checked in this figure, all quantities, linking numbers difference, excess of twist, and writhe, whether local or global, remain strictly constant along the trajectories. No relaxation actually ever occurs, and conformations obtained after long integration times are still composed of one torsionally relaxed moiety and one torsionally stressed moiety, as can be checked in Fig. 3 c. This third model of DNA-bridging proteins finally describes a topological barrier. To act as topological barriers, DNA-bridging proteins must consequently exert rather strong constraints on their binding sites. Indeed, they must block the diffusion of the excess of twist through both binding sites and must additionally block the rotation of one segment relative to the other one. It is worth emphasizing that this last condition is probably not satisfied by all DNA-bridging proteins. For example, it is well known that H-NS dimers bind most often nonspecifically to the DNA, which means that the two C-terminal DNA-binding domains (59) located at the end of flexible linkers (60) bind into the minor groove of the DNA (59) through generic electrostatic forces instead of a precise chemical bond. The consequence is that H-NS dimers can slide along the DNA molecule (28,29), but they are also expected to rotate with respect to the DNA axis. Conversely, the two DNA segments can rotate with respect to the protein axis. Proteins binding specifically to the DNA are more likely to satisfy the no-rotation condition, provided that the bonds are strong enough to resist the torque exerted by the DNA. A second mechanism for satisfying the no-rotation condition is that the proteins systematically form pairs of neighboring bridges. In this respect, we note...
with interest that the authors of the experimental study about the topological barrier formed by LacI tetramers reported that the barrier formed by a single LacI tetramer is much less stable compared with those containing multiple LacI-lac OI nucleoprotein complexes (15). The most straightforward explanation for this result is that two bridges in tandem are required to sustain the torque arising from the difference in torsional stress in the two DNA loops. Yet, single molecule experiments have demonstrated that a single LacI bridge can sustain the torque associated with physiological values of supercoiling, that is $\sigma \approx -0.06$ (16). In this context, we surmise that the slow relaxation observed in (15) for single DNA bridges may correspond to the slow intrication obtained with our second model for DNA-bridging proteins, whereas the more stable topological separation observed in (15) for bridges in tandem corresponds to true barriers obtained with the third model for DNA-bridging proteins. We finally note that the intrication properties discussed here are also expected to contribute to the formation of topological domains that are, in effect, larger than the domain delineated by the binding sites of LacI, as recently observed in (16).

**CONCLUSIONS**

DNA-bridging proteins form DNA loops by dynamically cross-linking DNA sites that are widely separated from the genomic point of view (17-22). They usually bind DNA nonspecifically but with high affinity. DNA-bridging proteins have now been investigated for more than two decades, but it is only very recently that it has been demonstrated that a DNA-bridging protein like the LacI repressor is able to separate circular plasmids into two topologically independent loops (15,23,24). To the best of our knowledge, this is, at present, the only demonstrated case of a DNA-bridging protein acting as a topological barrier. In this work, we aimed at going one step further and determining the conditions that DNA-bridging proteins must fulfill to act as topological barriers. We showed that these proteins must exert rather strong constraints on their binding sites: they must block the diffusion of the excess of twist through the two binding sites on the DNA molecule and must simultaneously prevent the rotation of one DNA segment relative to the other one. These criteria should be of great help when investigating the (existence, or lack of) topological barrier properties of other DNA-bridging proteins, like for example H-NS, which has been predicted to participate in the organization of the genome into topologically independent domains (6,61). Indeed, it is probable that not all DNA-bridging proteins satisfy the second condition. If it is not satisfied, then intricate conformations are obtained, with a preserved difference in the torsional stress between the two domains but different values of the stress in each domain. A rationale for the intriguing limit $W_{12} = Wr$ (or equivalently, $W_1 + W_2 = 0$) is missing, however.

This work also brings some light on another mechanism that has been proposed to explain the formation of independent topological domains, namely two proteins wrapping the DNA in two different locations ((15), Fig. 6). There is experimental evidence that certain wrapping proteins, like GalR (15), the $\lambda$ O protein (15), and Fis (62), are able to form topological barriers by blocking the diffusion of twist. However, we checked that the torsional stress equilibrates rapidly in the two moieties of the DNA chain when simulations are performed with $d_{\text{agg}} = 10$ nm instead of 4 nm because equilibration of twist and writhe is mediated in this case by the DNA chain passing repeatedly between beads $\alpha$ and $\beta$. The fact that GalR and $\lambda$ O proteins are able to separate the plasmid into two stable topological domains (15) therefore suggests that the two wrapping proteins are located very close to each other or are bound by some impenetrable molecular complex. Moreover, their relative orientation probably remains constant because the intrication mechanism would otherwise set in. Our work therefore clearly supports the hypothesis of the authors, who state that they cannot fully exclude the role of protein-protein interactions of $\lambda$-O-DNA complexes and GalR-DNA complexes in the formation of the two distinct topological domains (15).

Still, our work suggests that the conditions for topological separation may be met by molecular geometries that do not require that the boundaries of the topological domains be close to each other. For example, one can imagine a topological barrier formed of two different proteins, which both anchor the DNA molecule to the cell membrane. According to our work, the DNA segment located between the two anchoring proteins may be topologically independent, provided that each protein is able to block the diffusion of twist along the DNA molecule and that the proteins cannot rotate with respect to the membrane. If this latter condition is not satisfied, then some complex winding of the DNA around itself is likely to occur, like for DNA-bridging proteins.

Another model that is frequently proposed for the formation of topological barriers, namely actively transcribing RNAP (10-12), is more difficult to discuss in terms of the results described in this study because several timescales need be considered. Indeed, it is likely that the two parts of the DNA molecule located, respectively, upstream and downstream from the transcribing RNAP will tend to increase their intricacy to reduce the torsional stress induced by the blocking of the diffusion of twist at the RNAP. However, the RNAP is itself moving forward along the DNA so that the ability of transcribing RNAP to act as topological barriers probably depends on the relative speeds of the two mechanisms. More work is needed to clarify this point. Similarly, the case of DNA-bending nucleoid proteins, like HU (63), certainly deserves further investigations.

Finally, we note with interest that the intricate structure obtained with DNA-bridging proteins that block the diffusion of twist along the DNA but do not prevent the relative rotation of the DNA segments are highly compact. It turns
out that the mechanism leading to the formation of the bacterial nucleoid is a longstanding but still lively debated question (31,32,64–67). The point that has puzzled scientists for decades is that the volume of the unconstrained genomic DNA of bacteria in physiological solution, as estimated for example from the worm-like chain model (68), is approximately 1000 times larger than the volume of the cell, whereas the nucleoid often occupies only a small fraction of the cell (69). It would certainly be interesting to check whether the intricacy resulting from the alternation of torsionally stressed and torsionally relaxed DNA domains could contribute significantly to the compaction of the nucleoid.

**SUPPORTING MATERIAL**

Supporting Material can be found online at https://doi.org/10.1016/j.bpj.2020.08.004.

**AUTHOR CONTRIBUTIONS**

I.J. imagined the research project and performed preliminary simulations. M.J. developed the models, ran the simulations, and analyzed the results. M.J. and I.J. wrote the manuscript.

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**SUPPORTING CITATIONS**

References (70,71) appear in the Supporting Material.

**REFERENCES**

Supplemental Information

Requirements for DNA-Bridging Proteins to Act as Topological Barriers of the Bacterial Genome

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Requirements for DNA-bridging proteins to act as topological barriers of the bacterial genome

- Supporting Material –

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MODEL AND SIMULATIONS

Temperature $T$ is assumed to be 298 K throughout the study. Circular DNA molecules are modeled as circular chains of $n = 600$ beads with radius $a = 1.0$ nm separated at equilibrium by a distance $l_0 = 2.5$ nm. Two beads represent 15 DNA base pairs (bp) and chains of $n = 600$ beads (equivalent to 4500 bp) model the plasmids investigated in (1). The potential energy of the system, $E_{pot}$, consists of four terms

$$E_{pot} = V_s + V_b + V_t + V_e ,$$  \hspace{1cm} (S1)

where

$$V_s = \frac{\hbar}{2} \sum_{k=1}^{n} (l_k - l_0)^2$$ \hspace{1cm} (S2)

$$V_b = \frac{g}{2} \sum_{k=1}^{n} \theta_k^2$$ \hspace{1cm} (S3)

$$V_t = \frac{\tau}{2} \sum_{k=1}^{n} (\Phi_{k+1} - \Phi_k)^2$$ \hspace{1cm} (S4)

$$V_e = q^2 \sum_{k=1}^{n-2} \sum_{K=k+2}^{n} H(\|r_k - r_K\| - 2a)$$ \hspace{1cm} (S5)
describe the stretching, bending, torsional, and electrostatic energy of the DNA chain, respectively. In Eqs. (S2)-(S5), \( r_k \) denotes the position of bead \( k \) (with the convention that \( r_{n+k} = r_k \)), \( l_k = \| r_{k+1} - r_k \| \) the distance between two successive beads \( k \) and \( k+1 \), and 
\[
\theta_k = \arccos((r_{k+1} - r_k)(r_{k+2} - r_{k+1})/\left(\|r_{k+1} - r_k\| \|r_{k+2} - r_{k+1}\|\right))
\]
the angle formed by three successive beads \( k, k+1 \), and \( k+2 \).

The stretching energy \( V_s \) is a computational device without biological meaning, which is aimed at avoiding a rigid rod description. The stretching rigidity \( h \) was set to \( h = 100 k_B T/l_0^2 \) to insure that the variations of the distance between successive beads remain small enough (2).

In contrast, the bending rigidity \( g \) in the expression of \( V_b \) was obtained from the known persistence length of DNA, \( \xi = 50 \) nm, according to \( g = \xi k_B T/l_0 = 20 k_B T \).

The torsional contribution \( V_t \) was borrowed from Ref. (3) and torsion forces and momenta were computed as described therein. \( \Phi_{k+1} - \Phi_k \) denotes the rotation of the body-fixed frame \((u_k, f_k, v_k)\) around \( u_k \) between successive beads \( k \) and \( k+1 \), where 
\[
u_k = (r_{k+1} - r_k)/\|r_{k+1} - r_k\|
\]
the unit vector pointing from bead \( k \) to bead \( k+1 \). By convention, \( \Phi_{n+k} = \Phi_k \). The value of the torsional rigidity, \( \tau = 25 k_B T \), was obtained by imposing that the writhe contribution \( W_r \) accounts for approximately 70% of the linking number difference \( \Delta L_k \) at equilibrium (4), see Fig. S1. The values of the bending and torsional rigidities of the model are close to each other, which agrees with experimental findings (4).

The electrostatic energy of the DNA chain, \( V_e \), is expressed as a sum of repulsive Debye-Hückel terms with hard core. Function \( H(r) \) is defined according to
\[
H(r) = \frac{1}{4\pi \varepsilon r} \exp\left(-\frac{r}{r_D}\right),
\]
where \( \varepsilon = 80 \varepsilon_0 \) denotes the dielectric constant of the buffer and \( r_D = 1.07 \) nm the Debye length inside the buffer. This value of the Debye length corresponds to a concentration of monovalent salt of 100 mM, which is the value that is generally assumed for the cytoplasm of bacterial cells. \( q \) is the value of the electric charge, which is placed at the centre of each DNA bead.
\[ q = -\frac{\ell_B}{e} \approx -3.52 \bar{e}, \quad (S7) \]

where \( \bar{e} \) is the absolute charge of the electron and \( \ell_B = 0.7 \text{ nm} \) the Bjerrum length of water.

In Eq. (S7), \( \bar{e}/\ell_B \) is the net linear charge density along a DNA molecule immersed in a buffer with monovalent cations derived from Manning’s counterion condensation theory (5,6). Note that electrostatic interactions between nearest neighbours are not included in Eq. (S5) because it is considered that they are already accounted for in the stretching and bending terms.

The dynamics of the system was investigated by integrating numerically overdamped Langevin equations. Practically, the updated positions and torsion angles at time step \( i+1 \) are computed from the positions and torsion angles at time step \( i \) according to

\[ r_k^{(i+1)} = r_k^{(i)} + \frac{\Delta t}{6\pi \eta} f_k^{(i)} + \sqrt{\frac{2 k_BT \Delta t}{6\pi \eta a^2}} x_k^{(i)}, \quad (S8) \]

\[ \Phi_k^{(i+1)} = \Phi_k^{(i)} + \frac{\tau \Delta t}{4\pi \eta a^2 I_0} (\Phi_{k+1}^{(i)} - 2\Phi_k^{(i)} + \Phi_{k-1}^{(i)}) + \sqrt{\frac{2 k_BT \Delta t}{4\pi \eta a^2 I_0}} X_k^{(i)}, \quad (S9) \]

where \( f_k^{(i)} \) are vectors of inter-particle forces arising from the potential energy \( E_{\text{pot}} \), \( T = 298 \text{ K} \) is the temperature of the system, \( x_k^{(i)} \) and \( X_k^{(i)} \) are vectors of random numbers extracted from a Gaussian distribution of mean 0 and variance 1, \( \eta = 0.00089 \text{ Pa s} \) is the viscosity of the buffer at 298 K, and \( \Delta t = 10 \text{ ps} \) is the integration time step.

The excess of twist \( \Delta T_w \), the writhe \( W_r \), and the linking number difference \( \Delta L_k \) of the DNA chain were computed at regular time intervals according to (3)

\[ \Delta T_w = \frac{1}{2\pi} \sum_{k=1}^{n} \left( \Phi_{k+1} - \Phi_k \right) \quad (S10) \]

\[ W_r = \frac{1}{4\pi} \sum_{k=1}^{n} \sum_{K=1, K \neq k}^{n} \frac{\left( (r_{k+1} - r_k) \times (r_{k+1} - r_K) \right) \cdot (r_k - r_K)}{|r_k - r_K|^3} \quad (S11) \]

\[ \Delta L_k = \Delta T_w + W_r. \quad (S12) \]

The superhelical density \( \sigma \) was subsequently estimated from

\[ \sigma = \frac{\Delta L_k}{L_{k_0}}, \quad (S13) \]

where the linking number \( L_{k_0} = 7.5 n/10.5 \approx 429 \) is the ratio of the number of base pairs of the DNA chain and the mean number of base pairs per turn of the torsionally relaxed double helix. Fig. S1 shows the plot of \( \langle W_r / \Delta L_k \rangle \), the mean contribution of the writhe to the linking
number difference, as a function of $-\sigma$. This plot is in good agreement both with experimental results (4) and results obtained with another model (7).

Partial values of the excess of twist and the writhe were also computed for each of the two moieties separated by beads $\alpha$ ($1 \leq \alpha \leq n/2$) and $\beta = \alpha + n/2$ (each moiety comprises $n/2$ beads). Partial values of the excess of twist were simply obtained from

$$\Delta T_{W_1} = \frac{1}{2\pi} \sum_{k=\alpha}^{\beta-1} (\Phi_{k+1} - \Phi_k)$$  \hspace{1cm} (S14)

$$\Delta T_{W_2} = \frac{1}{2\pi} \sum_{k=\beta}^{\beta+n/2-1} (\Phi_{k+1} - \Phi_k).$$  \hspace{1cm} (S15)

The sum of $\Delta T_{W_1}$ and $\Delta T_{W_2}$ is obviously equal to the total excess of twist

$$\Delta T_W = \Delta T_{W_1} + \Delta T_{W_2}. \hspace{1cm} (S16)$$

The case of the writhe is more complex. Partial values were obtained by defining two closed chains of length $n/2$, namely $C_1 = (r_{\alpha}, r_{\alpha+1}, r_{\alpha+2}, ..., r_{\beta-2}, r_{\beta-1}, r_{\alpha})$ and $C_2 = (r_{\beta}, r_{\beta+1}, r_{\beta+2}, ..., r_{\alpha}, r_{\beta}, r_{\beta-1}, r_{\alpha})$, and computing the writhe for $C_1$ and $C_2$ (labeled $W_{T_1}$ and $W_{T_2}$, respectively) according to equations similar to Eq. (S11). Note that $C_1$ and $C_2$ are “virtual” chains, because torsional stress cannot flow directly between beads $\alpha$ and $\beta$.

Most importantly, $W_T$ is not necessarily close to the sum of $W_{T_1}$ and $W_{T_2}$, because of the double sums in the expressions for $W_T$, $W_{T_1}$, and $W_{T_2}$. Instead, one has

$$W_T = W_{T_1} + W_{T_2} + W_{T_{12}}. \hspace{1cm} (S17)$$

$W_T$, $W_{T_1}$, and $W_{T_2}$ quantify the winding of each closed chain around itself. In contrast, $W_{T_{12}}$ quantifies the winding of $C_1$ around $C_2$, that is, loosely speaking, the intrication of the two loops when beads $\alpha$ and $\beta$ remain close to each other.

Finally, the global torsional state of each moiety can be described by the value of the partial linking number difference

$$\Delta L_{k_1} = \Delta T_{W_1} + W_{T_1}$$  \hspace{1cm} (S18)

$$\Delta L_{k_2} = \Delta T_{W_2} + W_{T_2}. \hspace{1cm} (S19)$$

It results from Eqs. (S12) and (S16)-(S19) that

$$\Delta L_k = \Delta L_{k_1} + \Delta L_{k_2} + W_{T_{12}}. \hspace{1cm} (S20)$$
SUPPORTING REFERENCES


Figure S1: Evolution of $\langle W_r / \Delta L_k \rangle$, the mean contribution of the writhe to the linking number difference, as a function of $-\sigma$. 
Figure S2: Time evolution of (a) $\Delta T_{w_2} - \Delta T_{w_1}$, and (b) $W_{r_2} - W_{r_1}$, for an initial conformation with $\Delta L k^\theta = -26$ ($\sigma^\theta \approx -0.061$). The solid red lines show the results obtained upon averaging of $\Delta T_{w_2} - \Delta T_{w_1}$ and $W_{r_2} - W_{r_1}$ over 100 different trajectories starting from the same initial conformation. The dashed blue lines show the result of exponential fits against the red lines. The time windows used in the fits are $0 \leq t \leq 20$ μs for $\Delta T_{w_2} - \Delta T_{w_1}$ and $10 \leq t \leq 1000$ μs for $W_{r_2} - W_{r_1}$. 
Figure S3: Same as Fig. 2, except that rotational noise has been discarded from evolution equations Eq. (S9). Plots (a) and (b) show the evolution of $\tau_{T_w}$ and $\tau_{W_r}$ as a function of $-\sigma^0$, respectively. The dot-dashed lines are linear fits to the data and are provided as guidelines to the eyes and as a means of comparison with Figs. 2 and S4.
Figure S4: Same as Fig. S3, except that a bond has been added between beads $\alpha$ and $\beta$, as the crudest model for a DNA-bridging protein (see Eq. (3)). Plots (a) and (b) show the evolution of $\tau_{T_\nu}$ and $\tau_{W_{\nu}}$ as a function of $-\sigma^0$, respectively. The dot-dashed lines are linear fits to the data and are provided as guidelines to the eyes and as a means of comparison with Figs. 2 and S3.
Figure S5: Time evolution of $W_{r_1}$, $W_{r_2}$, and $W_{r_{12}}$ during relaxation of a torsionally out-of-equilibrium DNA conformation with $\Delta L k^0 = -52$ ($\sigma^0 \approx -0.121$) and (a) no constraint on the DNA chain, or (b) a bond between beads $\alpha$ and $\beta$ (Eq. (3)). Each plot was obtained by averaging $W_{r_1}$, $W_{r_2}$, and $W_{r_{12}}$ over 100 different trajectories starting from the same initial conformation.
Figure S6: Time evolution of $W_{12}$ during the relaxation of a torsionally out-of-equilibrium DNA conformation with $\Delta Lk^0 = -20 \ (\sigma^0 \approx -0.047)$ constrained by the second model for DNA-bridging proteins (Eqs. (5) and (6)). The four curves correspond to four different trajectories starting from the same initial conformation.
Figure S7: Time evolution of (a) the excess of twist, and (b) the writhe, during the relaxation of a torsionally out-of-equilibrium DNA conformation with \( \Delta L k^0 = -52 \) \( (\sigma^0 \approx -0.121) \) constrained by the third model for DNA-bridging proteins (Eqs. (5) and (7)). Each curve was averaged over 2 different trajectories starting from the same initial conformation.