SURVEY AND SUMMARY
Are TADs supercoiled?

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ABSTRACT
Topologically associating domains (TADs) are megabase-sized building blocks of interphase chromosomes in higher eukaryotes. TADs are chromosomal regions with increased frequency of internal interactions. On average a pair of loci separated by a given genomic distance contact each other 2–3 times more frequently when they are in the same TAD as compared to a pair of loci located in two neighbouring TADs. TADs are also functional blocks of chromosomes as enhancers and their cognate promoters are normally located in the same TAD, even if their genomic distance from each other can be as large as a megabase. The internal structure of TADs, causing their increased frequency of internal interactions, is not established yet. We survey here experimental studies investigating presence of supercoiling in interphase chromosomes. We also review numerical simulation studies testing whether transcription-induced supercoiling of chromatin fibres can explain how TADs are formed and how they can assure very efficient interactions between enhancers and their cognate promoters located in the same TAD.

INTRODUCTION
From topological domains to TADs

Many of us are familiar with the famous dictum attributed to Jacques Monod ‘Anything found to be true of Escherichia coli must also be true of elephants’ (1). That dictum nicely grasps the essence of the unifying principle of biology that applies to such fundamentals of life as DNA structure, genetic code, gene regulation, mechanisms of DNA replication etc. Of course, the details of a given process, like of DNA recombination, for example, can differ between E. coli and elephant, i.e. between prokaryotes and higher eukaryotes, but such general features like action of recombinase proteins that stretch and unwind the DNA during the homologous paring process are universal (2).

For >50 years now, it has been known that bacterial chromosomes with their several MB long circular DNA molecules are organized into topological domains, where sequential, 5–50 kb large portions of DNA form supercoiled loops (3–5). We progressively started to appreciate the significance of topological domains with respect to such aspects as giving bacterial chromosomes an organized structure (5) or providing the possibility of having different supercoiling level in different topological domains, which in turn permits a differential control of gene expression (6). Knowing the importance of topological domains for gene regulation in bacteria, it is natural to consider whether organization of chromosomes into supercoiled loops, that can vary their supercoiling level, belongs to fundamental biological principles that are maintained during evolution from prokaryotes to higher eukaryotes.

At least since 1977, when the well-known now electron microscopy pictures of histone-depleted metaphase chromosomes were published by Paulson and Laemmli (7), it has been considered as demonstrated that metaphase chromosomes of higher eukaryotes are composed of ca 100 kb long loops. The chromosome loops were also proposed to organize interphase chromosomes (8), where at least part of these loops involved matrix attachment regions interacting with nuclear lamin (9) and type II DNA topoisomerases bound to chromatin near loop attachment sites (10). These loops were frequently called topological domains (11), without directly implying though that they are supercoiled, as that is the case of bacterial topological domains. However, it was also shown that many of detected chromatin loops resulted from non-specific chromatin aggregation (12) and there were no chromatin loops revealed by cryo-electron microscopy studies of chromosomes (13). The lack of consensus in the field contributed to the fact that the concept of
organization of interphase chromosomes into specific loops gave stage to other developments in chromosome research that in the last two decades were connected to progress in genomics methods. The view that dominated the field and textbooks at the turn of 21st century was that during interphase stage of the cell cycle, chromatin fibres just behave like very long linear polymers that, due to their large size, require very long time to equilibrate (14–16). Only quite recently, though, thanks to very rapid development of high-resolution 3C (Chromosome Conformation Capture) methods, almost everybody in chromosome structure field started to think and talk about topological domains in interphase chromosomes of higher eukaryotes. In 2012, three seminal papers using advanced 3C methods revealed that human, mouse or drosophila chromosomes are composed of ca 1MB large, self-interacting regions that follow each other along individual chromosomes (17–19). On average, for the same genomic distance, two chromosomal loci contained in the same self-interacting domain contact each other 2–3 times more frequently than two loci located in two neighbouring domains. The title of one of those seminal papers even announced the identification of topological domains in mammalian genomes (17). The term ‘topological domains’ implicitly suggested that the structure of self-interacting domains in eukaryotic chromosomes is likely to be similar to the structure of topological domains in bacterial chromosomes. However, the model of self-interacting domains proposed in that seminal paper of Dixon et al. was very different from the structure of supercoiled topological domains in bacterial chromosomes. Individual topological domains in mammalian chromosomes were schematically presented as quasi-spherical globules (17). Dixon et al. did not discuss what underlying physical and biological mechanisms could be responsible for the formation of proposed chromatin globules with increased frequency of internal contacts. Supercoiling was not considered as a possible mechanism of compacting individual self-interacting domains (17). In another paper published in the same issue of Nature, Nora et al. mapped positions of self-interacting chromatin domains in a specific region of mice interphase X chromosome (18). Nora et al. named these self-interacting chromatin domains as topologically associating domains (TADs). Since then, the acronym TAD is very widely used and almost needs no explanation in today’s papers on chromosome structure.

Models of TADs

Probably the first model of TADs formation was proposed by Barbieri et al. (20). In their model, chromatin portions constituting different TADs were assumed to have different epigenetic marks, which in turn bind mark-specific polyvalent binders. Polyvalent binders compactify then individual TADs, which in turn increases the frequency of intra-TAD contacts. That model of TADs formation requires that neighbouring TADs should have different epigenetic marks, as otherwise such neighbouring TADs would fuse. However, at least in mammalian cells, large chromatin portions with the same epigenetic state (transcriptionally active, or inactive, for example) are still composed of individual TADs (17). Therefore, some other mechanism should be responsible for TADs formation in chromosomal portions with the same epigenetic state.

Modelling studies inspired by bacterial topological domains tested whether supercoiling of chromatin fibres can reproduce several known features of eukaryotic chromosomes (21). Models where individual TADs formed a few plectonemic superturns reproduced qualitatively the experimental contact maps (Figure 1). This included ca. 2–3-fold increase of intra-TAD contacts as compared to inter-TADs contacts, as well as the scaling relation between the probability of contacts and the genomic distance between interacting loci, which for genomic distances of the order of TAD sizes was characterized by the scaling exponent $\alpha$ of $-0.6$ (18).

Although models assuming supercoiling of chromatin fibres forming individual TADs were able to qualitatively reproduce the experimental data (21), the question remains whether TADs are indeed supercoiled and what are the mechanisms of supercoiling generation in eukaryotic chromosomes. Eukaryotes, in contrast to bacteria, do not have gyrase that is a specialized DNA topoisomerase that actively supercoils the DNA using the energy gained from ATP hydrolysis. However, transcription is known to be a potent generator of supercoiling both in prokaryotes and eukaryotes. Since late 1980ies it is known that transcribing RNA polymerases by binding to various nuclear components, including other RNA polymerases are prevented from encircling the transcribed template (22,23). As a consequence, transcribed DNA and thus transcribed chromatin fibres are forced to undergo axial rotation (24–28). The axial rotation of chromatin fibres is generated at sites where helical DNA is actually passing through RNA polymerase. The transmission of rotational motion to flanking portions of chromatin fibres is however slow in viscous nuclear environment that limits diffusion of torsional stress along chromatin fibres (29) and even a friction between DNA and interior of type I topoisomerases limits swivelling motion needed for the relaxation of the torsional stress (30). As a consequence, negative supercoiling accumulates behind transcribing polymerases, whereas ahead of transcribing polymerases one observes formation of positive supercoiling (24–27). Cellular DNA topoisomerases can relax both positive and negative supercoiling. However, the efficiency of relaxation of positive and negative supercoiling are not the same (31,32). Positive supercoiling is relaxed more rapidly as its accumulation is more harmful for the cells by its inhibitive effect on the process of DNA replication and transcription (32). Therefore, ongoing transcription and action of cellular topoisomerases can maintain complex dynamic equilibrium of DNA supercoiling where different portions of chromosomes can have different levels of DNA supercoiling. Numerous studies over the last 30 years provided fragmentary view on supercoiling of specific portions of the genome (33–35). Only during the last decade, with the development of modern genomic approaches, we gained insights into the global supercoiling landscape of eukaryotic chromosomes (31,36–38). A convenient method to study DNA supercoiling is based on the quantification of the level of binding of intercalating agents such as psoralen to DNA (31). Binding of intercalating drugs decreases torsional stress in negatively super-
coiled DNA, whereas it increases the torsional stress in covalently closed non-supercoiled DNA or in positively supercoiled DNA. Therefore, when psoralen is added at low concentration, the probability of its binding and crosslink formation increases with the magnitude of negative supercoiling of a given topological domain, where it can be detected and quantified upon UV crosslinking (31). Differential levels of psoralen binding to various genomic regions can be detected using such genomic approaches as high-resolution microarrays (36). Using this approach, Naughton et al. have provided the evidence that human interphase chromosomes are divided into domains with various levels of supercoiling (36). Actively transcribed regions showed high affinity to psoralen leading to a conclusion that they are negatively supercoiled. The transcriptionally inactive regions showed diminished affinity to psoralen, which is consistent with the idea that these regions are non-supercoiled. Diminished level of psoralen binding to transcriptionally inactive chromatin regions could be also caused by positive supercoiling resulting from formation of additional nucleosomes, as each formed nucleosome introduces one positive supercoil (39). Transcriptionally inactive regions may also be bound by heterochromatin binding proteins that make DNA less accessible to psoralen.

Both, transcription and topoisomerase activities were shown to be required to maintain the steady state dynamic profile of supercoiling with chromosomal portions showing various levels of psoralen binding. The fact that neighbouring chromatin domains frequently show very different affinity to psoralen indicated that these domains form chromatin loops with different levels of supercoiling and these loops are separated by borders that do not permit the supercoiling to diffuse between neighbouring loops (36). In addition, Naughton et al., reported that the borders between chromatin regions with different levels of DNA supercoiling frequently corresponded to inter TADs borders detected by 3C methods (36). However, on average, chromatin domains with a quasi-constant level of DNA supercoiling had the median size of ca 100 kb, which is ca 10 times smaller than the average size of TADs determined in the early studies (17) but is a typical size of smaller loops within TADs known as sub-TADs (40). It would be very informative to compare high-resolution Hi-C-determined borders of TADs and sub-TADS with the borders of supercoiling domains determined by Naughton et al. (36).

Using a similar genomic approach based on preferential binding of psoralen to negatively supercoiled DNA, Kouzine et al. mapped more precisely where transcription-induced supercoiling localizes (37). Meta-analysis of all regions surrounding transcription start sites (TSS) of active genes in human cells indicated that transcription-generated negative supercoiling was strongest near TSSs (37). Traces of transcription-induced positive supercoiling were detected ahead of transcribing polymerases. However, the magnitude of positive supercoiling was much smaller than the magnitude of negative supercoiling detected near TSS (37). This later result indicates that positive supercoiling is relaxed much quicker by cellular topoisomerases than negative supercoiling. Similar conclusion was also reached in earlier studies of transcription-induced supercoiling in yeast (32). Also, recent studies of transcription in human

Figure 1. Models of supercoiled TADs show consistency with experimental data. (A) Experimental contact map of a short chromosome region with two TADs. That map was generated using the deposited data (18) and shows two TADs that were indicated with letters E and F in the contact map shown in Figure 1c of Nora et al. paper (18). (B) Contact maps obtained in a simulation of two neighbouring chromatin loops that were modelled as supercoiled elastic polymers with properties approximating physical behaviour of chromatin fibres with 400 and 800 kb, respectively. In this simulation the supercoiling density was set to two negative superturns for every 100 kb. (C) Comparisons between the experimental and simulated contact frequency profiles. Experimental data for the genomic region shown in (A) are from Nora et al. (18). The experimental results are shown as individual data points, whereas simulated results form continuous profiles. The straight dashed line shows the slope, which would correspond to the rate of contact decrease characterized by the exponent of ~0.6. Intra-TAD contacts determined experimentally or by simulations are indicated with one letter description of the corresponding TADs. Two letter descriptions indicate corresponding inter-TAD contacts. Red and blue colour relate to intra- and inter-TAD contacts, respectively. Notice that simulations give directly the contact probability whereas experimental data give only relative contact probabilities. Therefore, on log/log plots, the experimental data points can be all together shifted vertically by the same rescaling factor during the fitting procedure. Figure 1 is adapted from (21).
Favourable environment for further gene expression (45). On ing in TADs containing active genes is expected to create as already discussed (see Figure 1), supercoiling of individual TADs would compactify them and therefore would increase the frequency of internal contacts. Therefore, supercoiling of individual TADs would provide a physical explanation of why, on average, a pair of loci located in the same TAD contact each other more frequently than a pair of loci separated by the same genomic distance but located in neighbouring TADs. This increase of the average intra-TAD contacts is of course important for enhancer-promoter contacts and explains in part the fact that enhancers and their cognate promoters are located in the same TAD (46). However, knowing how critical is the proper enhancer-promoter interaction for cellular differentiation and ontogenesis of complex organisms, such as humans (46), one realizes that the predicted 2–3-fold difference in the contact probability between legitimate (cognate) and illegitimate (non-cognate) enhancer-promoter pairs is unlikely to be sufficient to guarantee a robust, error-free development of multicellular organisms. The 2–3-fold preference for cognate contact would result in non-cognate contact for two or three cognate contacts, which would result in ectopic transcription of many genes and frequent lack of transcription of genes that should be active in a particular setting. Let us therefore reanalyse what we know about the effect of supercoiling on the contacts of genetic loci located in the same supercoiled loop.

Vologodskii et al., in their early simulation studies of supercoiled DNA molecules, arrived to the conclusion that supercoiling dramatically increases the frequency of intramolecular contacts between DNA sites that are far from each other along the contour of the same supercoiled DNA molecule (47). This supercoiling-induced increase of DNA-DNA contact probability was estimated to be of about two orders of magnitude as compared to non-supercoiled DNA molecules (47). These 1992 results are widely quoted and it is now a very popular notion that supercoiling dramatically increases frequency of contacts between sites located in the same supercoiled DNA molecule. However, if supercoiling were indeed increasing the frequency of intramolecular contacts by two orders of magnitude it would speak against the proposal that TADs are constituted by supercoiled loops. The frequencies of intra- and inter-TAD contacts between neighbouring TADs should then also differ by two orders of magnitude, whereas in reality there is only a 2–3-fold enhancement of intra TAD contacts as detected by 3C methods. Only in 2001, new simulation studies by Vologodskii et al. (48) brought a new important aspect into light. When they correctly took into account that under physiological conditions the electrostatic repulsion between DNA segments is nearly completely screened (49), the stimulation of contacts by supercoiling was greatly diminished (48). In their 1992 simulation studies Vologodskii et al. considered the effect of supercoiling on DNA maintained under low salt conditions i.e. conditions of strong electrostatic repulsion between DNA segments (39,49). Under these conditions supercoiling acts against electrostatic repulsion and stimulates DNA-DNA contacts that only occur very rarely in non-supercoiled DNA. Therefore, the two orders of magnitude increase of intra-molecular contacts by DNA supercoiling only applies to physiologically irrelevant regime where DNA segments strongly repulse each other. Under physiological conditions electrostatic charges of DNA and chromatin are practically neutralized which makes that DNA-DNA contacts are frequent also without

Facilitation of local separation of DNA strands by negative supercoiling

Early studies addressing the role of negative supercoiling in bacteria revealed that negative supercoiling, which is known to facilitate local strand separation, stimulates the process of transcription (42). The facilitation of local strand separation by negative supercoiling should also apply to eukaryotic transcription and numerous experiments showed that this is indeed the case (43,44). Thus negative supercoiling in TADs containing active genes is expected to create favourable environment for further gene expression (45). On the other hand, positive supercoiling or just the absence of negative supercoiling make the strand separation more difficult, which in turn would inhibit initiation of transcription. In this respect the observation of Naughton et al. (36) that negatively supercoiled chromosomal domains contain active genes, whereas positively or less negatively supercoiled domains group inactive genes suggests that supercoiling by its direct effect on the stability of DNA may be used to regulate gene expression in eukaryotic cells.

Supercoiling stimulates legitimate enhancer-promoter contacts

As already discussed (see Figure 1), supercoiling of individual TADs would compactify them and therefore would increase the frequency of internal contacts. Therefore, supercoiling of individual TADs would provide a physical
supercoiling and therefore the expected stimulation of intra-TAD contacts by supercoiling becomes very modest (21).

However, this modest, 2–3-fold stimulation of contacts brings us back to the initial problem that such a low stimulation of intra-TAD contacts may not be sufficient to ensure that enhancers act only on their cognate promoters located in the same TAD and are not involved in ectopic activation of genes in the neighbouring TADs. There is, however, a possible solution to this problem. In previous simulation studies investigating the frequency of contacts in supercoiled DNA molecules or chromatin fibres the authors investigated only the contacts between generic portions of modelled DNA or chromatin fibres, where the contacting portions had no mutual affinities (21,47,48). Enhancers and promoters, though, show mutual affinity mediated by specific transcription factors such as mediator or YY1, for example (50). Thus, the relevant question is how the interactions between sites that show mutual affinity are stimulated by supercoiling under ionic conditions corresponding to physiological conditions where electrostatic repulsion between DNA segments or chromatin fibres is neutralized. To answer this question Benedetti et al. proceeded with Brownian dynamics simulations of supercoiled DNA molecules that either were composed only of generic beads that were not attracting each other or contained in addition two distally located beads that were showing a short-range attraction to each other (51). This mutual attraction had the strength and other characteristics mimicking protein-mediated DNA–DNA interactions. The simulations revealed that in modelled DNA molecules containing only generic sites with no mutual affinity to each other the fraction of time during which a pair of pre-assigned distally located sites stayed in contact with each other was practically not increased by supercoiling (see Figure 2A). However, simulated DNA molecules containing two sites with mutual affinity reacted to increasing supercoiling by several dozen-fold increase of the fraction of time these sites with mutual affinity remained in a contact (Figure 2A). To understand better why supercoiling stimulate so strongly interactions of sites with mutual affinities Benedetti et al. analysed the on and off times of modelled enhancer-promoter interactions. Interestingly, the on times of enhancer-promoter interaction i.e. the average duration of the bound state before it undergoes thermally induced dissociation, was practically the same for supercoiled and non-supercoiled systems. However, the off time i.e. the average time separating the moments of dissociation from the moments of re-binding dramatically decreased as supercoiling was introduced. Shortening of the average off time by supercoiling indicates that after enhancer-promoter dissociation the two partners were kept close to each other. This is the consequence of the fact that supercoiling naturally keeps interwound segments in close proximity to each other and a complex and slow slithering motion is needed to move formerly interacting sites further from each other. Therefore, in supercoiled systems there is a high chance that after thermally induced dissociation of enhancer-promoter complex this complex can reform again. In case of non-supercoiled systems or in case of ectopic contacts between enhancer located in two different supercoiled TADs after thermal dissociation of such an enhancer-promoter interaction the two

Figure 2. Supercoiling stimulates intramolecular interactions involving sites with mutual affinity. (A) Supercoiling increases the fraction of time during which a pair of sites with mutual affinity stays together, whereas interactions of sites with no mutual affinity is hardly affected by supercoiling. Simulations were set to mimic physical behaviour of 3 kb long circular DNA molecules, however qualitatively similar results are also expected for much larger, closed chromatin loops. The sites whose mutual contacts were followed, were placed 180° apart on the circular map of the modelled DNA molecules. The strength of the short-range interaction between the two sites with mutual affinity was set to 10 kBT (ca. 6 kcal/mol), which is in the intermediate range of strengths of protein-DNA interactions (89). The blue profile shows how the fraction of time during which two sites with mutual affinities stay together increases with increasing supercoiling. Without supercoiling that fraction of time is close to 0 since two sites with the affinity of 10 kBT easily dissociate due to thermal fluctuations and then in non-supercoiled DNA molecules it takes very long time till they form a new contact. The red profile shows the fraction of time during which the two sites with no mutual affinity stay together as supercoiling increases. Simulation snapshots show molecules with two sites with mutual affinity (blue) or with no mutual affinity (red). The corresponding supercoiling level is indicated. The ΔLk = −15 indicates that the linking number of the modelled DNA molecule is diminished by 15 turns as compared to torsionally relaxed DNA molecules of the same size. 3 kb long DNA molecules with ΔLk = −15 have their density of supercoiling corresponding to this of deproteinized plasmid DNA molecules isolated from healthy bacterial cells. (B) Supercoiling diminishes the average duration of the OFF state involving two sites with mutual affinities. The inset shows the telegraphic profile registering duration of ON and OFF states. The ON state is characterized by the distance between centres of beads representing enhancer and promoter being smaller than 6 nm. Otherwise the two beads are in the OFF state. Notice that increasing supercoiling decreases the duration of the average OFF state by over two orders of magnitude (blue profile). Figure 2 is adapted from (51).
partners can quickly diffuse from each other, which limits their chances of quick rebinding.

Benedetti et al. simulated also the situation where an enhancer had the choice to interact with the promoter located in the same TAD or in the neighbouring TAD (see Figure 3). When both TADs were not supercoiled, the fraction of time the enhancer spent in intradomain (legitimate) interactions was ca 3 times larger than the fraction of time spent in interdomain (illegitimate) interactions. However, when both TADs were supercoiled, the intradomain (legitimate) interactions were ca. 30-fold favoured over interdomain (illegitimate) interactions. Therefore, in a biological setting where enhancers show affinity to promoters and where electrostatic repulsion between chromatin regions is largely screened, the supercoiling of chromosomal loops forming TADs can guarantee high fidelity of enhancer promoter interactions even if on average the contacts between generic sites located in the same TAD are only 2–3 times more frequent than the contacts between generic sites located in neighbouring TADs.

**Why is it advantageous to maintain large genomic distances between enhancers and their cognate promoters?**

Promoters of housekeeping genes, which are constitutively expressed in every tissue, do not require activating contacts with distally located enhancers but just need interaction with their proximal regulatory elements. Due to their mutual proximity promoters and regulatory elements of housekeeping genes can find each other very efficiently. Promoters of developmentally regulated genes, however, whose expression is induced only in certain tissues or only during certain periods of development, require for their activation an interaction with distally located enhancers (52, 53). Enhancers and their partner promoters are almost always located in the same TAD (46), they are frequently separated by large genomic distances which may exceed 1 MB but on average are of ~200 kb (54). It is relatively easy to understand that regulation of genes that should be only active in certain tissues or during certain period of development may be quite complex and it is very important that these genes will not be expressed at any other time than needed. Expression of developmentally regulated genes requires co-presence of specific transcription factors where some of them bind to enhancer and others bind to promoters. Only when the entire set of transcription factors specific for a given developmental stage or tissue is present, a given set of developmentally regulated genes can be activated by interactions involving direct contact between cognate enhancers and promoters (55). However, it is difficult to understand why it is advantageous to have enhancers located up to a megabase apart from their cognate promoters. To answer this question Benedetti et al., performed simulations of TADs in which the effect of supercoiling was investigated on enhancer-promoter pairs placed at different genomic distances from each other (51). They have observed that when the interacting sites with mutual affinities were placed at small genomic distances from each other they very efficiently interacted with each other irrespective of whether modelled TADs were supercoiled or not (51). However, when the interacting sites with mutual affinities were placed at large genomic distances from each other the supercoiling was needed to observe their efficient binding to each other (51). This result suggests that during evolution large genomic distances between enhancers and promoters were favoured, as only in this case the interaction between enhancers and promoters can be regulated by varying the level of supercoiling. Only distal location of enhancers with respect to promoters provides the possibility of this additional level of positive and negative control of developmentally regulated genes. Therefore, the fact that enhancers are usually located at large genomic distance from their cognate promoters supports the notion that supercoiling of chromatin can be used to regulate gene expression of genes in higher eukaryotes. This would be then similar to the role of supercoiling in gene regulation in bacteria where entire portions of chromosome can be transcriptionally activated or deactivated by changes of supercoiling level (56–58) or where transcriptional bursting depends on gyrase activity (59).
Transcription-induced supercoiling and chromatin loop extrusion

While all TADs can be recognized on Hi-C contact maps as ‘triangles’ delineating the regions with increased frequency of internal contacts, in approximately 50% of cases the formed triangles have very strong apaxes. TADs with such contact patterns form chromatin loops with their border elements contacting each other (60). ChIP-Seq studies revealed that contacting chromatin regions at the base of formed loops are bound by CTCF protein and cohesin (60). Interestingly, loop forming TADs are bordered specifically by CTCF binding sites in convergent orientation (60,61). A question of how long chromatin loops can be spanned only by contacts between CTCF proteins in convergent orientation have led to a model of chromatin loop extrusion (61–63). According to that model, individual or double rings of cohesin assembly on chromatin in such a way that a small loop is formed. That loop was proposed to grow then as cohesin rings slide along the embraced chromatin fibres (61–63). The movement of cohesin rings was proposed to stop after cohesin binds to C-terminal part of CTCF proteins and only convergent orientation of CTCF binding sites could expose C-terminal part of both CTCF proteins for stabilizing contacts with approaching cohesin rings (64). Numerical simulations testing such a mechanism closely reproduced experimental contact maps (61,62). However, until now it is not known what can drive chromatin loop extrusion in interphase chromosomes. Numerous experimental tests did not detect any DNA or chromatin translocation activity of cohesin. DNA translocation activity has been observed though in case of condensin (65), which forms a related protein complex to cohesin but which acts during condensation of mitotic or meiotic chromosomes. It is therefore possible that also cohesin has such an activity but it requires some specific conditions not found yet (66). However, it is also possible that cohesin is a DNA/chromatin translocase and needs to be pushed along chromatin fibres by other motor proteins such as RNA polymerase (67) or by other mechanisms (68).

Recent experiments by Vian et al. (63) showed that cohesin that is unable to hydrolyse ATP, due to a mutation in ATPase motif, is still capable of diffusing along chromatin and reaching CTCF at TADs borders, although this occurs up to 2-fold less efficiently as compared to not-mutated cohesin. The authors interpreted this diminished efficiency as an indication that cohesin is a motor protein that requires ATP for its translocation along the chromatin, although they considered also the possibility that ATPase mutations decrease cohesin loading on chromatin fibres and this in turn decreases that amount of cohesin that reaches CTCF anchors (63). It seems though that the second possibility is more likely in light of studies showing that cohesin ATPase activity is required for cohesin loading on chromatin (69). Therefore, the experimental observation that cohesin that is unable to hydrolyse ATP and thus loads less efficiently still reaches its final destination in >50% of cases (63), suggests that it is not ATPase activity of cohesin that is implicated in its translocation but some other factors, such as transcription-induced supercoiling, for example.

In vitro experiments aimed to study diffusional motion of cohesin rings along DNA and chromatin established that cohesin rings embrace chromatin fibres very tightly, which results in a strong molecular friction between cohesin and enclosed chromatin fibre (70). That friction greatly limits spontaneous diffusion of cohesin rings along chromatin but does not block it completely (70). The same study concluded that it is unlikely that individual cohesin rings can embrace two chromatin fibres and therefore two cohesin rings forming so called cohesin handcuffs would be needed to span the growing loop during chromatin loop extrusion (70).

Another development important for the understanding of the possible involvement of supercoiling in chromatin loop extrusion consisted of establishing that type II DNA topoisomerases are bound to CTCF proteins forming borders of loop-forming TADs (71,72). This location of type II DNA topoisomerases suggests that torsional stress generated during transcription within individual TADs can be dissipated by diffusing towards TADs borders. Studies of Baranello et al. on the other hand explained better how the interplay between transcribing RNA polymerases and interacting with them type I DNA topoisomerases results in the preferential relaxation of positive supercoiling generated by transcribing RNA polymerase and thus in a net input of negative supercoiling by ongoing transcription (41).

Racko et al., incorporated the mentioned above elements into simulation studies testing whether transcription-induced supercoiling could drive chromatin loop extrusion (73). In the modelled system, handcuffs-forming cohesin rings were placed on chromatin fibres in such a way that a small chromatin loop formed between the two rings. It was assumed then that transcription starts in that small loop. This assumption agrees with studies showing that cohesin rings are preferentially loaded by Nibbl protein in the immediate vicinity of transcription start sites (TSS) of active genes and their enhancers (67,74). Instead of modelling RNA polymerase producing domains with positive and negative supercoiling and then modelling DNA topoisomerase relaxing preferentially positive supercoiling, Racko et al. in their simulations, introduced directly negative supercoiling into the chromatin portion between the two cohesin rings. In principle, when positive supercoils generated during transcription are preferentially relaxed by DNA topoisomerases, transcription is able to introduce one negative supercoil for every 10 nucleotides that are transcribed (75). This negative supercoiling is generated and initially confined to the chromatin portion behind transcribing RNA polymerase (24). Since transcribing RNA polymerase blocks the diffusion of supercoiling through the template portion it is actually bound to (24), dissociation of RNA polymerase would be needed to permit diffusion and equilibration of negative supercoiling in the entire chromatin loop between two cohesin rings. However, it is known that transcription cycles in well over 90% of cases undergo premature termination with RNA polymerase dissociating after synthesis of short 30–75 nt long RNAs (76,77). Racko et al., therefore assumed that RNA polymerase dissociation permits the negative supercoiling accumulated behind transcribing RNA polymerase to diffuse and equilibrate in the entire chromatin portion between two cohesin rings (73). It was also assumed that this process of injecting and equili-
brating of negative supercoiling in the chromatin portion between two cohesin rings occurs repeatedly (73).

As cohesins rings embrace tightly chromatin fibres and this slows down motion of chromatin fibres passing through cohesin rings (70) Racko et al. in their simulations increased hydrodynamic drag of regions of the chain that actually pass through modelled cohesin handcuffs. Accounting for this effect limits the dissipation of supercoiling via diffusion through cohesin rings forming the handcuff and thus favours accumulation of supercoiling in chromatin loops spanned by cohesin handcuffs (See Figure 4). Accounting for experimental demonstration that CTCF proteins defining boundaries of TADs are associated with type II DNA topoisomerases (71), Racko et al. introduced into modelled chromatin fibres short regions where segments can freely pass through each other and where they can also freely swivel (see Figure 4).

Figure 4 shows snapshots from simulations testing whether transcription-induced supercoiling can drive chromatin loop extrusion (73). Transcription occurring in the chromatin portion spanned by cohesin handcuffs causes formation of a plectoneme. As more supercoiling is introduced, the formed plectoneme grows and pushes the cohesin handcuffs towards the borders of the modelled TAD. That process can be also described as pulling of chromatin fibres through cohesin rings. The movement of chromatin fibres with respect to cohesin handcuffs is formally analogous to the process of chromatin loop extrusion proposed earlier (61,62). However, previous models assumed that cohesin has DNA/chromatin translocation activity or is pushed by some translocases, whereas simulations by Racko et al. showed that transcription-induced supercoiling is capable of driving chromatin loop extrusion (73).

Very recent studies by Vian et al. (63) concluded that chromatin loop extrusion can serve as a reeling mechanism ensuring that active enhancers progressively scan chromatin regions separating them from their cognate promoters. For this to happen the two cohesin rings forming a handcuff should be placed initially on both sides of an enhancer TSS and then one of the rings should be immobilized by its stabilizing interaction with CTCF located close to the enhancer. Progressing chromatin loop extrusion permits then the enhancer to scan for stabilizing contacts within up to megabase large stretches of chromatin loops forming individual TADs. An analogous mechanism, but based on supercoiling-driven movement of chromatin fibres with respect of cohesin handcuffs, was proposed earlier (73) and is presented here in Figure 5.

It is important to stress here that the proposal involving supercoiling in formation of TADs critically depends on the presence of cohesin and is therefore entirely consistent with all studies showing the importance of cohesin in subdividing interphase chromosomes into TADs with CTCF-defined border elements (63,78–80).

There is growing evidence that transcription of enhancers and formation of enhancer promoter contacts precedes the transcription of genes activated by these enhancers (81,82) as it is also shown in the model in Figure 5. However, in the majority of cases the produced eRNAs does not seem to have a specific role and are quickly degraded (83). This has led to proposals that the mere act of tran-
A significant confusion regarding the role of transcription in chromatin-loop extrusion may be caused by the fact that TADs in Drosophila were shown to form independently of transcription (85). However, TADs in Drosophila might not be formed by chromatin loop extrusion and/or that other mechanisms seem to play an important role (86). TADs in Drosophila are nanocompartments (87) i.e. short continuous chromatin portions with the same types of epigenetic marks. Direct or protein-mediated attraction between the same types of epigenetic marks causes then an increased frequency of internal contacts in these nanocompartments (87). The ‘hallmarks’ of the loop extrusion process, which are clearly visible in Hi-C contact maps of mammalian cells are focal peaks resulting from prolonged direct contacts between two borders of a given TAD (60,61).

In mammalian cells, nearly half of TADs show such focal peaks (60) whereas in Drosophila cells such peaks are visible only in <1% of regions with increased frequency of contacts (86). As clearly stated by Eagen et al., ‘A close correlation between TADs and loops detectable by Hi-C is not conserved throughout metazoans.’ (86).

**CONCLUSIONS**

We discussed here experimental data supporting the notion that TADs are supercoiled. We also presented how supercoiling of chromatin fibres can be generated during transcription and how supercoiling is expected to induce compaction of individual TADs and thus promote interaction between enhancers and their cognate promoters that are located in the same TAD. In addition, we propose that transcription-induced supercoiling is likely to provide a long-sought solution to the question of how chromatin

**Figure 5.** Schematic drawing explaining how an asymmetric loop extrusion starting from an enhancer located near CTCF sites drives chromatin scanning by that enhancer. (A) Cohesin rings preferentially load around transcribed enhancers (67,74). eRNA transcription, combined with the relaxation of positive supercoiling by Top1 (41,84), injects net negative supercoiling into the portion of chromatin fibre that is flanked by two cohesin rings. Due to the accumulated supercoiling, the enhancer region is drawn into a plectoneme and stays in a close contact with a stretch of chromatin that actually faces it. (B, C) While short eRNAs are produced, more negative supercoiling is injected causing the plectoneme to grow. Since one of cohesin rings is ‘clogged’ by its interaction with CTCF, the formed plectoneme can only grow and thus decrease its torsional stress by pulling chromat in fibre through the other cohesin ring forming the handcuff. The growth of plectoneme requires internal slithering motion that is well studied in case of supercoiled DNA (48). The portions of chromatin pulled through un clogged cohesin ring slide along the enhancer region. This sliding motion, or scanning, is the basis of a progressive 1D search during which enhancers with bound transcription factors can very efficiently find their cognate promoters to which they show affinity. Figure 5 is adapted from (73).
loop extrusion is driven and explains the mechanical role of eRNA transcription. New experimental approaches, such as single cell Hi-C (88) are needed, however, to reveal to us the structure of individual TADs. It is important to add here that transcription-induced supercoiling would be only needed to stimulate the expression of genes positively regulated by contacts with distal regulatory elements. Transcription of eRNA or of house-keeping genes which do not need an activation by interaction with distal regulatory elements can be initiated in the absence of supercoiling and can be used then as the source of supercoiling activating other genes.

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